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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 95/18858		
C12N 15/19, C07K 14/52, 16/24 38/19	A61K A1	(43) International Publication Date: 13 July 1995 (13.07.95)		
(21) International Application Number:  (22) International Filing Date: 28 Dece  (30) Priority Data:  08/176,553 3 January 1994 08/185,607 21 January 199 08/196,689 15 February 19 08/223,263 4 April 1994 (( 08/249,376 25 May 1994 ( 08/348,657 2 December 19 08/348,658 2 December 19	4 (21.01.94) U 94 (15.02.94) U 4.04.94) U 25.05.94) U	(71) Applicant (for all designated States except US): GENENTECH.		
US Filed on 4 US Filed on 15 Feb US Filed on 2 Dece US Filed on 21 Ja US Filed on 2 Dece US Filed on 22 Dece US Filed on 22 Dece	08/249,376 (CII May 1994 (25.05.9- 08/223,263 (CII April 1994 (04.04.9- 08/196,689 (CII ruary 1994 (15.02.9- 08/348,658 (CII mber 1994 (02.12.9- 08/185,607 (CII nuary 1994 (21.01.9- 08/348,657 (CII mber 1994 (02.12.9- 08/176,553 (CII nuary 1994 (03.01.9-	CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TJ, TT, UA, US, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD, SZ).  Published  With international search report.  Before the expiration of the time limit for amending the		

## (54) Title: THROMBOPOIETIN

#### (57) Abstract

Isolated thrombopoietin (TPO), isolated DNA encoding TPO, and recombinant or synthetic methods of preparing and purifying TPO are disclosed. Various forms of TPO are shown to influence the replication, differentiation or maturation of blood cells, especially megakaryocytes and megakaryocyte progenitor cells. Accordingly, these compounds may be used for treatment of thrombocytopenia.

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# **THROMBOPOIETIN**

#### FIELD OF THE INVENTION

This invention relates to the isolation, purification and recombinant or chemical synthesis of proteins that influence survival, proliferation, differentiation or maturation of hematopoletic cells, especially platelet progenitor cells. This invention specifically relates to the cloning and expression of nucleic acids encoding a protein ligand capable of binding to and activating *mpl*, a member of the cytokine receptor superfamily. This invention further relates to the use of these proteins alone or in combination with other cytokines to treat immune or hematopoletic disorders including thrombocytopenia.

#### **BACKGROUND OF THE INVENTION**

#### I. The Hematopoietic System

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The hematopoietic system produces the mature highly specialized blood cells known to be necessary for survival of all mammals. These mature cells include; erythrocytes, specialized to transport oxygen and carbon dioxide, T- and B-lymphocytes, responsible for cell- and antibody-mediated immune responses, platelets or thrombocytes, specialized to form blood clots, and granulocytes and macrophages, specialized as scavengers and as accessory cells to combat infection. Granulocytes are further subdivided into; neutrophils, eosinophils, basophils and mast cells, specialized cell types having discrete functions. Remarkably, all of these specialized mature blood cells are derived from a single common primitive cell type, referred to as the pluripotent (or totipotent) stem cell, found primarily in bone marrow (Dexter et al., Ann. Rev. Cell Biol., 3:423-441 [1987]).

The mature highly specialized blood cells must be produced in large numbers continuously throughout the life of a mammal. The vast majority of these specialized blood cells are destined to remain functionally active for only a few hours to weeks (Cronkite *et al.*, *Blood Cells*, 2:263-284 [1976]). Thus, continuous renewal of the mature blood cells, the primitive stem cells themselves, as well as any intermediate or lineage-committed progenitor cell lines lying between the primitive and mature cells, is necessary in order to maintain the normal steady state blood cell needs of the mammal.

At the heart of the hematopoietic system lies the pluripotent stem cell(s). These cells are relatively few in number and undergo self-renewal by proliferation to produce daughter stem cells or are transformed, in a series of differentiation steps,

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into increasingly mature lineage-restricted progenitor cells, ultimately forming the highly specialized mature blood cell(s).

For example, certain multipotent progenitor cells, referred to as CFC-Mix, derived from stem cells undergo proliferation (self-renewal) and development to produce colonies containing all the different myeloid cells; erythrocytes, neutrophils, megakaryocytes (predecessors of platelets), macrophages, basophils, eosinophils, and mast cells. Other progenitor cells of the lymphoid lineage undergo proliferation and development into T-cells and B-cells.

Additionally, between the CFC-Mix progenitor cells and myeloid cells lie another rank of progenitor cells of intermediate commitment to their progeny. These lineage-restricted progenitor cells are classified on the basis of the progeny they produce. Thus, the known immediate predecessors of the myeloid cells are: erythroid colony-forming units (CFU-E) for erythrocytes, granulocyte/macrophage colony-forming cells (GM-CFC) for neutrophils and macrophages, megakaryocyte colony-forming cells (Meg-CFC) for megakaryocytes, eosinophil colony-forming cells (Eos-CFC) for eosinophils, and basophil colony-forming cells (Bas-CFC) for mast cells. Other intermediate predecessor cells between the pluripotent stem cells and mature blood cells are known (see below) or will likely be discovered having varying degrees of lineage-restriction and self-renewal capacity.

The underlying principal of the normal hematopoietic cell system appears to be decreased capacity for self-renewal as multipotency is lost and lineage-restriction and maturity is acquired. Thus, at one end of the hematopoietic cell spectrum lies the pluripotent stem cell possessing the capacity for self-renewal and differentiation into all the various lineage-specific committed progenitor cells. This capacity is the basis of bone marrow transplant therapy where primitive stem cells repopulate the entire hematopoietic cell system. At the other end of the spectrum lie the highly lineage-restricted progenitors and their progeny which have lost the ability of self-renewal but have acquired mature functional activity.

The proliferation and development of stem cells and lineage-restricted progenitor cells is carefully controlled by a variety of hematopoietic growth factors or cytokines. The role of these growth factors in vivo is complex and incompletely understood. Some growth factors, such as interleukin-3 (IL-3), are capable of stimulating both multipotent stem cells as well as committed progenitor cells of several lineages, including for example, megakaryocytes. Other factors such as granulocyte/macrophage colony-stimulating factor (GM-CSF) was initially thought to be restricted in its action to GM-CFC's. Later, however, it was discovered GM-CSF also influenced the proliferation and development of interalia megakaryocytes. Thus, IL-3 and GM-CSF were found to have overlapping biological activities, although with

differing potency. More recently, both interleukin-6 (IL-6) and interleukin-11 (IL-11), while having no apparent influence on meg-colony formation alone, act synergistically with IL-3 to stimulate maturation of megakaryocytes (Yonemura et al., Exp. Hematol., 20:1011-1016 [1992]).

Thus, hematopoletic growth factors may influence growth and differentiation of one or more lineages, may overlap with other growth factors in affecting a single progenitor cell line, or may act synergistically with other factors.

It also appears that hematopoietic growth factors can exhibit their effect at different stages of cell development from the totipotent stem cell through various committed lineage-restricted progenitors to the mature blood cell. For example, erythropoietin (epo) appears to promote proliferation only of mature erythroid progenitor cells. IL-3 appears to exert its effect earlier influencing primitive stem cells and intermediate lineage-restricted progenitor cells. Other growth factors such as stem cell factor (SCF) may influence even more primitive cell development.

It will be appreciated from the foregoing that novel hematopoietic growth factors that affect survival, proliferation, differentiation or maturation of any of the blood cells or predecessors thereof would be useful, especially to assist in the reestablishment of a diminished hematopoietic system caused by disease or after radiation- or chemo-therapy.

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### II. Megakaryocytopoiesis - Platelet Production

Regulation of megakaryocytopoiesis and platelet production has been reviewed by: Mazur, Exp. Hematol., 15:248 [1987] and Hoffman, Blood, 74:1196-1212 Briefly, bone marrow pluripotent stem cells differentiate into [1989]. megakaryocytic, erythrocytic, and myelocytic cell lines. It is believed there is a hierarchy of committed megakaryocytic progenitor cells between stem cells and megakaryocytes. At least three classes of megakaryocytic progenitor cells have been identified, namely; burst forming unit megakaryocytes (BFU-MK), colony-forming unit megakaryocytes (CFU-MK), and light density megakaryocyte progenitor cells (LD-CFU-MK). Megakaryocytic maturation itself is a continuum of development that has been separated into stages based on standard morphologic criteria. The earliest recognizable member of the megakaryocyte (MK or meg) family are the megakaryoblasts. These cells are initially 20 to 30 μm in diameter having basophilic cytoplasm and a slightly irregular nucleus with loose, somewhat reticular chromatin and several nucleoli. Later, megakaryoblasts may contain up to 32 nuclei (ployploid). but the cytoplasm remains sparse and immature. As maturation proceeds, the nucleus becomes more lobulate and pyknotic, the cytoplasm increases in quantity and becomes more acidophilic and granular. The most mature cells of this family may give the

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appearance of releasing platelets at their periphery. Normally, less than 10% of megakaryocytes are in the blast stage and more than 50% are mature. Arbitrary morphologic classifications commonly applied to the megakaryocyte series are megakaryoblast for the earliest form; promegakaryocyte or basophilic megakaryocyte for the intermediate form; and mature (acidophilic, granular, or platelet-producing) megakaryocyte for the late forms. The mature megakaryocyte extends filaments of cytoplasm into sinusoidal spaces where they detach and fragment into individual platelets (Williams et al., Hematology, 1972).

Megakaryocytopoiesis is believed to involve several regulatory factors (Williams et al., Br. J. Haematol., 52:173 [1982] and Williams et al., J. Cell Physiol., 110:101 [1982]). The early level of megakaryocytopoiesis is postulated as being mitotic, concerned with cell proliferation and colony initiation from CFU-MK but is not affected by platelet count (Burstein et al., J. Cell Physiol., 109:333 [1981] and Kimura et al., Exp. Hematol., 13:1048 [1985]). The later stage of maturation is non-mitotic, involved with nuclear polyploidization and cytoplasmic maturation and is probably regulated in a feedback mechanism by peripheral platelet number (Odell et al., Blood, 48:765 [1976] and Ebbe et al., Blood, 32:787 [1968]).

The existence of a distinct and specific megakaryocyte colony-stimulating factor (MK-CSF) has been disputed (Mazur, *Exp. Hematol.*, 15:340-350 [1987]). However most authors believe that a process so vital to survival as platelet production would be regulated by cytokine(s) exclusively responsible for this process. The hypothesis that megakaryocyte/platelet specific cytokine(s) exist has provided the basis for more than 30 years of search - but to date no such cytokine has been purified, sequenced and established by assay as a unique MK-CSF (TPO).

Although it has been reported that MK-CSF's have been partly purified from experimentally produced thrombocytopenia (Hill et al., Exp. Hematol., 14:752 [1986]) and human embryonic kidney conditioned medium [CM] (McDonald et al., J. Lab. Clin. Med., 85:59 [1975]) and in man from a plastic anemia and idiopathic thrombocytopenic purpura urinary extracts (Kawakita et al., Blood, 6:556 [1983]) and plasma (Hoffman et al., J. Clin. Invest., 75:1174 [1985]), their physiological function is as yet unknown in most cases.

The conditioned medium of pokeweed mitogen-activated spleen cells (PWM-SpCM) and the murine myelomonocyte cell line WEHI-3 (WEHI-3CM) have been used as megakaryocyte potentiators. PWM-SpCM contains factors enhancing CFU-MK growth (Metcalf et al., Pro. Natl. Acad. Sci., USA, 72:1744-1748 [1975]; Quesenberry et al., Blood, 65:214 [1985]; and Iscove, N.N., in Hematopoietic Cell Differentiation, ICN-UCLA Symposia on Molecular and Cellular Biology, Vol. 10, Golde

et al., eds. [New York, Academy Press] pp 37-52 [1978]), one of which is interleukin-3 (IL-3), a multilineage colony stimulating factor (multi-CSF [Burstein, Blood Cells, 11:469 [1986]). The other factors in this medium have not yet been identified and isolated. WEHI-3 is a murine myelomonocytic cell line secreting relatively large amounts of IL-3 and smaller amounts of GM-CSF. IL-3 has been found to potentiate the growth of a wide range of hematopoietic cells (Ihle et al., J. Immunol., 13:282 [1983]). IL-3 has also been found to synergize with many of the known hematopoietic hormones or growth factors (Bartelmez et al., J. Cell Physiol., 122:362-369 [1985] and Warren et al., Cell, 46:667-674 [1988]), including both erythropoletin (EPO) and interleukin-1 (IL-1), in the induction of very early multipotential precursors and the formation of very large mixed hematopoietic colonies.

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Other sources of megakaryocyte potentiators have been found in the conditioned media of murine lung, bone, macrophage cell lines, peritoneal exudate cells and human embryonic kidney cells. Despite certain conflicting data (Mazur, Exp. Hematol., 15:340- 350 [1987]), there is some evidence (Geissler et al., Br. J. Haematol., 60:233-238 [1985]) that activated T lymphocytes rather than monocytes play an enhancing role in megakaryocytopoiesis. These findings suggest that activated Tlymphocyte secretions such as interleukins may be regulatory factors in MK development (Geissler et al., Exp. Hematol., 15:845-853 [1987]). A number of studies on megakaryocytopolesis with purified erythropoletin EPO (Valnchenker et al., Blood, 54:940 [1979]; McLeod et al., Nature, 261:492-4 [1976]; and Williams et al., Exp. Hematol., 12:734 [1984]) indicate that this hormone has an enhancing effect on MK colony formation. This has also been demonstrated in both serum-free and serum-containing cultures and in the absence of accessory cells (Williams et al., Exp. Hematol., 12:734 [1984]). EPO was postulated to be involved more in the single and two-cell stage aspects of megakaryocytopoiesis as opposed to the effect of PWM-SpCM which was involved in the four-cell stage of megakaryocyte development. The interaction of all these factors on both early and late phases of megakaryocyte development remains to be elucidated.

Data produced from several laboratories suggests that the only multi-lineage factors that individually have MK-colony stimulating activity are GM-CSF and IL-3 and, to a lesser extent, the B-cell stimulating factor IL-6 (Ikebuchi et al., Proc. Natl. Acad. Sci. USA, 84:9035 [1987]). More recently, several authors have reported that IL-11 and leukemia inhibitory factor (LIF) act synergistically with IL-3 to increase megakaryocyte size and ploidy (Yonemura et al., British Journal of Hematology, 84:16-23 [1993]; Burstein et al., J. Cell. Physiol., 153:305-312 [1992]; Metcalf et al., Blood, 76:50-56 [1990]; Metcalf et al., Blood, 77:2150-2153 [1991];

Bruno et al., Exp. Hematol., 19:378-381 [1991]; and Yonemura et al., Exp. Hematol., 20:1011-1016 [1992]).

Other documents of interest include: Eppstein et al., U.S. Patent No. 4,962,091; Chong, U.S. Patent No. 4,879,111; Fernandes et al., U.S. Patent No. 4,604,377; Wissler et al., U.S. Patent No. 4,512,971; Gottlieb, U.S. Patent No. 4,468,379; Bennett et al., U.S. Patent No. 5,215,895; Kogan et al., U.S. Patent No. 5,250,732; Kimura et al., Eur. J. Immunol., 20(9):1927-1931 [1990]; Secor et al., J. of Immunol., 144(4):1484-1489 [1990]; Warren et al., J. of Immunol., 140(1):94-99 [1988]; Warren et al., Exp. Hematol., 17(11):1095-1099 [1989]; Bruno et al., Exp. Hematol., 17(10):1038-1043 [1989]; Tanikawa et al., Exp. Hematol., 17(8):883-888 [1989]; Koike et al., Blood, 75(12):2286-2291 [1990]; Lotem, Blood, 75(5):1545-1551 [1989]; Rennick et al., Blood, 73(7):1828-1835 [1989]; and Clutterbuck et al., Blood, 73(6):1504-1512 [1989].

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## III. Thrombocytopenia

Platelets are critical elements of the blood clotting mechanism. Depletion of the circulating level of platelets, called thrombocytopenia, occurs in various clinical conditions and disorders. Thrombocytopenia is commonly defined as a platelet count below 150 X 10<sup>9</sup> per liter. The major causes of thrombocytopenia can be broadly divided into three categories on the basis of platelet life span, namely; (1) impaired production of platelets by the bone marrow, (2) platelet sequestration in the spleen (splenomegaly), or (3) increased destruction of platelets in the peripheral circulation (e.g., autoimmune thrombocytopenia or chemo- and radiation-therapy). Additionally, in patients receiving large volumes of rapidly administered platelet-poor blood products, thrombocytopenia may develop due to dilution.

The clinical bleeding manifestations of thrombocytopenia depend on the severity of thrombocytopenia, its cause, and possible associated coagulation defects. In general, patients with platelet counts between 20 and 100 X 10<sup>9</sup> per liter are at risk of excessive post traumatic bleeding, while those with platelet counts below 20 X 10<sup>9</sup> per liter may bleed spontaneously. These latter patients are candidates for platelet transfusion with attendant immune and viral risk. For any given degree of thrombocytopenia, bleeding tends to be more severe when the cause is decreased production rather than increased destruction of platelets. In the latter situation, accelerated platelet turnover results in the circulation of younger, larger and hemostatically more effective platelets. Thrombocytopenia may result from a variety of disorders briefly described below. A more detailed description may be found in Schafner, A. I., "Thrombocytopenia and Disorders of Platelet Function," *Internal* 

Medicine, 3rd Ed., John J. Hutton et al., Eds., Little Brown and Co., Boston/Toronto/London [1990].

(a) Thrombocytopenia due to impaired platelet production

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Causes of congenital thrombocytopenia include constitutional aplastic anemia (Fanconi syndrome) and congenital amegakaryocytic thrombocytopenia, which may be associated with skeletal malformations. Acquired disorders of platelet production are caused by either hypoplasia of megakaryocytes or ineffective thrombopoiesis. Megakaryocytic hypoplasia can result from a variety of conditions, including marrow aplasia (including idiopathic forms or myelosuppression by chemotherapeutic agents or radiation therapy), myelfibrosis, leukemia, and invasion of the bone marrow by metastatic tumor or granulomas. In some situations, toxins, infectious agents, or drugs may interfere with thrombopoiesis relatively selectively; examples include transient thrombocytopenias caused by alcohol and certain viral infections and mild thrombocytopenia associated with the administration of thiazide diuretics. Finally, ineffective thrombopoiesis secondary to megaloblastic processes (folate or B12 deficiency) can also cause thrombocytopenia, usually with coexisting anemia and leukopenia.

Current treatment of thrombocytopenias due to decreased platelet production depends on identification and reversal of the underlying cause of the bone marrow failure. Platelet transfusions are usually reserved for patients with serious bleeding complications, or for coverage during surgical procedures, since isoimmunization may lead to refractoriness to further platelet transfusions. Mucosal bleeding resulting from severe thrombocytopenia may be ameliorated by the oral or intravenous administration of the antifibrinolytic agents. Thrombotic complications may develop, however, if antifibrinolytic agents are used in patients with disseminated intravascular coagulation (DIC).

## (b) Thrombocytopenia due to splenic sequestration

Splenomegaly due to any cause may be associated with mild to moderate thrombocytopenia. This is a largely passive process (hypersplenism) of splenic platelet sequestration, in contrast to the active destruction of platelets by the spleen in cases of immunomediated thrombocytopenia discussed below. Although the most common cause of hypersplenism is congestive splenomegaly from portal hypertension due to alcoholic cirrhosis, other forms of congestive, infiltrative, or lymphoproliferative splenomegaly are also associated with thrombocytopenia. Platelet counts generally do not fall below 50 X 10<sup>9</sup> per liter as a result of hypersplenism alone.

## (c) Thrombocytopenia due to nonimmune-mediated platelet destruction

Thrombocytopenia can result from the accelerated destruction of platelets by various nonimmunologic processes. Disorders of this type include disseminated intravascular coagulation, prosthetic intravascular devices, extra corporeal circulation of the blood, and thrombotic microangiopathies such as thrombotic thrombocytic purpura. In all of these situations, circulating platelets that are exposed to either artificial surfaces or abnormal vascular intima either are consumed at these sites or are damaged and then prematurely cleared by the reticuloendothelial system. Disease states or disorders in which disseminated intravascular coagulation (DIC) may arise are set forth in greater detail in Braunwald et al. (eds), Harrison's Principles of Internal Medicine, 11th Ed., p.1478, McGraw Hill [1987]. Intravascular prosthetic devices, including cardiac valves and intra-aortic balloons can cause a mild to moderate destructive thrombocytopenia and transient thrombocytopenia in patients undergoing cardiopulmonary bypass or hemodialysis may result from consumption or damage of platelets in the extra corporeal circuit.

#### (d) Drug-induced immune thrombocytopenia

More than 100 drugs have been implicated in immunologically mediated thrombocytopenia. However, only quinidine, quinine, gold, sulfonamides, cephalothin, and heparin have been well characterized. Drug-induced thrombocytopenia is frequently very severe and typically occurs precipitously within days while patients are taking the sensitizing medication.

#### (e) Immune (autoimmune) thrombocytopenic purpura (ITP)

ITP in adults is a chronic disease characterized by autoimmune platelet destruction. The autoantibody is usually IgG although other immunoglobulins have also been reported. Although the autoantibody of ITP has been found to be associated with platelet membrane GPII<sub>D</sub>III<sub>a</sub>, the platelet antigen specificity has not been identified in most cases. Extravascular destruction of sensitized platelets occurs in the reticuloendothelial system of the spleen and liver. Although over one-half of all cases of ITP are idiopathic, many patients have underlying rheumatic or autoimmune diseases (e.g., systemic lupus erythematosus) or lymphoproliferative disorders (e.g., chronic lymphocytic leukemia).

#### (f) HIV-Induced ITP

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ITP is an increasingly common complication of HIV infection (Morris et al., Ann. Intern. Med., 96:714-717 [1982]), and can occur at any stage of the disease progression, both in patients diagnosed with the Acquired Immune Deficiency Syndrome (AIDS), those with AIDS-related complex, and those with HIV infection but without AIDS symptoms. HIV infection is a transmissible disease ultimately characterized by a profound deficiency of cellular immune function as well as the

occurrence of opportunistic infection and malignancy. The primary immunologic abnormality resulting from infection by HIV is the progressive depletion and functional impairment of T lymphocytes expressing the CD4 cell surface glycoprotein (Lane et al., Ann. Rev. Immunol., 3:477 [1985]). The loss of CD4 helper/inducer T cell function probably underlies the profound defects in cellular and humoral immunity leading to the opportunistic infections and malignancies characteristic of AIDS (H. Lane supra).

Although the mechanism of HIV-associated ITP is unknown, it is believed to be different from the mechanism of ITP not associated with HIV infection. (Walsh et al., N. Eng. J. Med., 311:635-639 [1984]; and Ratner, Am. J. Med., 86:194-198 [1989]).

### IV. Current Therapy for Thrombocytopenia

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The therapeutic approach to the treatment of patients with thrombocytopenia is dictated by the severity and urgency of the clinical situation. The treatment is similar for HIV-associated and non-HIV-related thrombocytopenia, and although a number of different therapeutic approaches have been used, the therapy remains controversial.

Platelet counts in patients diagnosed with thrombocytopenia have been successfully increased by glucocorticoid (e.g., prednisolone) therapy, however in most patients, the response is incomplete, or relapse occurs when the glucocorticoid dose is reduced or its administration is discontinued. Based upon studies with patients having HIV-associated ITP, some investigators have suggested that glucocorticoid therapy may result in predisposition to AIDS. Glucocorticoids are usually administered if platelet count falls below 20 X 109/liter or when spontaneous bleeding occurs.

For patients refractory to glucocorticoids, the compound:

4-(2-chlorphenyl)-9-methyl-2-[3-(4-morpholinyl)-3-propanon-1-yl]6H-thieno[3,2,f][1,2,4]triazolo[4,3,a,][1,4]diazepin (WEB 2086)

has been successfully used to treat a severe case of non HIV-associated ITP. A patient having platelet counts of 37,000-58,000/µI was treated with WEB 2086 and after 1-2 weeks treatment platelet counts increased to 140,000-190,000/µI. (EP 361,077 and Lohman *et al.*, Lancet, 1147 [1988]).

Although the optimal treatment for acquired amegakaryocytic thrombocytopenia purpura (AATP) is uncertain, antithymocyte globulin (ATG), a horse antiserum to human thymus tissue, has been shown to produce prolonged complete remission (Trimble *et al.*, *Am. J. Hematol.*, 37:126-127 [1991]). A recent report however, indicates that the hematopoietic effects of ATG are attributable to thimerosal, where presumably the protein acts as a mercury carrier (Panella *et al.*, *Cancer Research*, 50:4429-4435 [1990]).

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Good results have been reported with splenectomy. Splenectomy removes the major site of platelet destruction and a major source of autoantibody production in many patients. This procedure results in prolonged treatment-free remissions in a large number of patients. However, since surgical procedures are generally to be avoided in immune compromised patients, splenectomy is recommended only in severe cases of thrombocytopenia (e.g. severe HIV-associated ITP), in patients who fail to respond to 2 to 3 weeks of glucocorticoid treatment, or do not achieve sustained response after discontinuation of glucocorticoid administration. Based upon current scientific knowledge, it is unclear whether splenectomy predisposes patients to AIDS.

In addition to prednisolone therapy and splenectomy, certain cytotoxic agents, e.g., vincristine, and azidothimidine (AZT, zidovudine) also show promise in treating HIV-induced ITP; however, the results are preliminary.

It will be appreciated from the foregoing that one way to treat thrombocytopenia would be to obtain an agent capable of accelerating the differentiation and maturation of megakaryocytes or precursors thereof into the platelet-producing form. Considerable efforts have been expended on identifying such an agent, commonly referred to as "thrombopoietin" (TPO). Other names for TPO commonly found in the literature include; thrombocytopoiesis stimulating factor (TSF), megakaryocyte colony-stimulating factor (MK-CSF), megakaryocytestimulating factor and megakaryocyte potentiator. TPO activity was observed as early as 1959 (Rak et al., Med. Exp., 1:125) and attempts to characterize and purify this agent have continued to the present day. While reports of partial purification of TPOactive polypeptides exist (see, for example, Tayrien et al., J. Biol. Chem., 262:3262 [1987] and Hoffman et al., J. Clin. Invest. 75:1174 [1985]), others have postulated that TPO is not a discrete entity in its own right but rather is simply the polyfunctional manifestation of a known hormone (IL- 3, Sparrow et al., Prog. Clin. Biol. Res., 215:123 [1986]). Regardless of its form or origin, a molecule possessing thrombopoietic activity would be of significant therapeutic value. Although no protein has been unambiguously identified as TPO, considerable interest surrounds the recent discovery that mpl, a putative cytokine receptor, may transduce a thrombopoletic signal.

## V. Mpl is a Megakaryocytopoletic Cytokine Receptor

It is believed that the proliferation and maturation of hematopoietic cells is tightly regulated by factors that positively or negatively modulate pluripotential stem cell proliferation and multilineage differentiation. These effects are mediated through the high-affinity binding of extracellular protein factors to specific cell surface receptors. These cell surface receptors share considerable homology and are generally

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classified as members of the cytokine receptor superfamily. Members of the superfamily include receptors for: IL-2 (β and γ chains) (Hatakeyama et al., Science, 244:551-556 [1989]; Takeshita et al., Science, 257:379-382 [1991]); IL-3 (Itoh et al., Science, 247:324-328 [1990]; Gorman et al., Proc. Natl. Acad. Sci. USA, 87:5459-5463 [1990]; Kitamura et al., Cell, 66:1165-1174 [1991a]; Kitamura et al., Proc. Natl. Acad. Sci. USA, 88:5082-5086 [1991b]), IL-4 (Mosley et al., Cell, 59:335-348 [1989], IL-5 (Takaki et al., EMBO J., 9:4367-4374 [1990]; Tavernier et al., Cell, 66:1175-1184 [1991]), IL-6 (Yamasaki et al., Science, 241:825-828 [1988]; Hibi et al., Cell, 63:1149-1157 [1990]), IL-7 (Goodwin et al., Cell, 60:941-951 [1990]), IL-9 (Renault et al., Proc. Natl. Acad. Sci. USA, 89:5690-5694 [1992]), granulocyte-macrophage colony-stimulating factor (GM-CSF) (Gearing et al., EMBO J., 8:3667-3676 [1991]; Hayashida et al., Proc. Natl. Acad. Sci. USA, 244:9655-9659 [1990]), granulocyte colony-stimulating factor (G-CSF) (Fukunaga et al., Cell, 61:341-350 [1990a]; Fukunaga et al., Proc. Natl. Acad. Sci. USA, 87:8702-8706 [1990b]; Larsen et al., J. Exp. Med., 172:1559-1570 [1990]), EPO (D'Andrea et al., Cell, 57:277-285 [1989]; Jones et al., Blood, 76:31-35 [1990]), Leukemia inhibitory factor (LIF) (Gearing et al., EMBO J., 10:2839-2848 [1991]), oncostatin M (OSM) (Rose et al., Proc. Natl. Acad. Sci. USA, 88:8641-8645 [1991]) and also receptors for prolactin (Boutin et al., Proc. Natl. Acad. Sci. USA, 88:7744-7748 [1988]; Edery et al., Proc. Natl. Acad. Sci. USA, 86:2112-2116 [1989]), growth hormone (GH) (Leung et al., Nature, 330:537-543 [1987]) and ciliary neurotrophic factor (CNTF) (Davis et al., Science, **253**:59-63 [1991].

Members of the cytokine receptor superfamily may be grouped into three functional categories (for review see Nicola *et al.*, *Cell*, **67**:1-4 [1991]). The first class comprises single chain receptors, such as erythropoletin receptor (EPO-R) or granulocyte colony stimulating factor receptor (G-CSF-R), which bind ligand with high affinity via the extracellular domain and also generate an intracellular signal. A second class of receptors, so called  $\alpha$ -subunits, includes interleukin-6 receptor (IL6-R), granulocyte-macrophage colony stimulating factor receptor (GM-CSF-R), interleukin-3 receptor (IL3-R $\alpha$ ) and other members of the cytokine receptor superfamily. These  $\alpha$ -subunits bind ligand with low affinity but cannot transduce an intracellular signal. A high affinity receptor capable of signaling is generated by a heterodimer between an  $\alpha$ -subunit and a member of a third class of cytokine receptors, termed  $\beta$ -subunits, *e.g.*,  $\beta_C$ , the common  $\beta$ -subunit for the three  $\alpha$ -subunits IL3-R $\alpha$  and GM-CSF-R.

Evidence that *mpl* is a member of the cytokine receptor superfamily comes from sequence homology (Gearing, *EMBO J.*, **8**:3667-3676 [1988]; Bazan, *Proc*.

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Natl. Acad. Sci. USA, 87:6834-6938 [1990]; Davis et al., Science, 253:59-63 [1991] and Vigon et al., Proc. Natl. Acad. Sci. USA, 89:5640-5644 [1992]) and its ability to transduce proliferative signals.

Deduced protein sequence from molecular cloning of murine c-mpl reveals this protein is homologous to other cytokine receptors. The extracellular domain contains 465 amino acid residues and is composed of two subdomains each with four highly conserved cysteines and a particular motif in the N-terminal subdomain and in the C-terminal subdomain. The ligand-binding extracellular domains are predicted to have similar double β-barrel fold structural geometries. This duplicated extracellular domain is highly homologous to the signal transducing chain common to IL-3, IL-5 and GM-CSF receptors as well as the low-affinity binding domain of LIF (Vigon et al., Oncogene, 8:2607-2615 [1993]). Thus mpl may belong to the low affinity ligand binding class of cytokine receptors.

A comparison of murine *mpl* and mature human *mpl* P, reveals these two proteins show 81% sequence identity. More specifically, the N-terminus and C-terminus extracellular subdomains share 75% and 80% sequence identity respectively. The most conserved *mpl* region is the cytoplasmic domain showing 91% amino acid identity, with a sequence of 37 residues near the transmembrane domain being identical in both species. Accordingly, *mpl* is reported to be one of the most conserved members of the cytokine receptor superfamily (Vigon *supra*).

Evidence that mpl is a functional receptor capable of transducing a proliferative signal comes from construction of chimeric receptors containing an extracellular domain from a cytokine receptor having high affinity for a known cytokine with the mpl cytoplasmic domain. Since no known ligand for mpl has been reported, it was necessary to construct the chimeric high affinity ligand binding extracellular domain from a class one cytokine receptor such as IL-4R or G-CSFR. Vigon et al., supra fused the extracellular domain of G-CSFR with both the transmembrane and cytoplasmic domain of c-mpl. An IL-3 dependent cell line, BAF/B03 (Ba/F3) was transfected with the G-CSFR/mpl chimera along with a full length G-CSFR control. Cells transfected with the chimera grew equally well in the presence of cytokine IL-3 or G-CSF. Similarly, cells transfected with G-CSFR also grew well in either IL-3 or G-CSF. All cells died in the absence of growth factors. A similar experiment was conducted by Skoda et al., EMBO J., 12(7):2645-2653 [1993] in which both the extracellular and transmembrane domains of human IL-4 receptor (hIL-4-R) were fused to the murine mpl cytoplasmic domain, and transfected into a murine IL-3 dependent Ba/F3 cell line. Ba/F3 cells transfected with wild type hIL-4-R proliferated normally in the presence of either of the species specific IL-4 or IL-3. Ba/F3 cells transfected with hIL-4R/mpl proliferated

normally in the presence of hIL-4 (in the presence or absence of IL-3) demonstrating that in Ba/F3 cells the *mpl* cytoplasmic domain contains all the elements necessary to transduce a proliferative signal.

These chimeric experiments demonstrate the proliferation signaling capability of the mpl cytoplasmic domain but are silent regarding whether the mpl extracellular domain can bind a ligand. These results are consistent with at least two possibilities, namely, mpl is a single chain (class one) receptor like EPO-R or G-CSFR or it is a signal transducing  $\beta$ -subunit (class three) requiring an  $\alpha$ -subunit like IL-3 (Skoda et al. supra).

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## VI. Mpl Ligand is a Thrombopoletin (TPO)

As described above, it has been suggested that serum contains a unique factor, sometimes referred to as thrombopoletin (TPO), that acts synergistically with various other cytokines to promote growth and maturation of megakaryocytes. No such natural factor has ever been isolated from serum or any other source even though considerable effort has been expended by numerous groups. Even though it is not known whether *mpl* is capable of directly binding a megakaryocyte stimulating factor, recent experiments demonstrate that *mpl* is involved in proliferative signal transduction from a factor or factors found in the serum of patients with aplastic bone marrow (Methia *et al.*, *Blood*, **82**(5):1395-1401 [1993]).

Evidence that a unique serum colony-forming factor distinct from IL-1 $\alpha$ , IL-3, IL-4, IL-6, IL-11, SCF, EPO, G-CSF, and GM-CSF transduces a proliferative signal through mpl comes from examination of the distribution of c-mpl expression in primitive and committed hematopoietic cell lines and from mpl antisense studies in one of these cell lines.

Using reverse transcriptase (RT)-PCR in immuno-purified human hematopoietic cells, Methia et al., supra demonstrated that strong mpl mRNA messages were only found in CD34+ purified cells, megakaryocytes and platelets. CD34+ cells purified from bone marrow (BM) represents about 1% of all BM cells and are enriched in primitive and committed progenitors of all lineages (e.g., erythroid, granulomacrophage, and megakaryocytic).

Mpl antisense oligodeoxynucleotides were shown to suppress megakaryocytic colony formation from the pluripotent CD34+ cells cultured in serum from patients with aplastic marrow (a rich source of megakaryocyte colony-stimulating activity [MK-CSA]). These same antisense oligodeoxynucleotides had no effect on erythroid or granulomacrophage colony formation.

Whether mpl directly bound a ligand and whether the serum factor shown to cause megakaryocytopolesis acted through mpl was still unknown. It had been

suggested, however, that if *mpl* did directly bind a ligand, its amino acid sequence was likely to be highly conserved and have species cross-reactivity owing to the considerable sequence identity between human and murine *mpl* extracellular domains (Vigon *et al.*, *supra* [1993]).

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#### VII. Objects

In view of the foregoing, it will be appreciated there is a current and continuing need in the art to isolate and identify molecules capable of stimulating proliferation, differentiation and maturation of hematopoietic cells, especially megakaryocytes or their predecessors for therapeutic use in the treatment of thrombocytopenia. It is believed such a molecule is a *mpl* ligand and thus there exists a further need to isolate such ligand(s) to evaluate their role(s) in cell growth and differentiation.

Accordingly, it is an object of this invention to obtain a pharmaceutically pure molecule capable of stimulating proliferation, differentiation and/or maturation of megakaryocytes into the mature platelet-producing form.

It is another object to provide the molecule in a form for therapeutic use in the treatment of a hematopoietic disorder, especially thrombocytopenia.

It is a further object of the present invention to isolate, purify and specifically identify protein ligands capable of binding *in vivo* a cytokine superfamily receptor known as *mpl* and to transduce a proliferative signal.

It is still another object to provide nucleic acid molecules encoding such protein ligands and to use these nucleic acid molecules to produce *mpl* binding ligands in recombinant cell culture for diagnostic and therapeutic use.

It is yet another object to provide derivatives and modified forms of the protein ligands including amino acid sequence variants, variant glycoprotein forms and covalent derivatives thereof.

It is an additional object to provide fusion polypeptide forms combining a *mpl* ligand and a heterologous protein and covalent derivatives thereof.

It is still an additional object to provide variant polypeptide forms combining a *mpl* ligand with amino acid additions and substitutions from the EPO sequence to produce a protein capable of regulating proliferation and growth of both platelets and red blood cell progenitors.

It is yet an additional object to prepare immunogens for raising antibodies against *mpl* ligands or fusion forms thereof, as well as to obtain antibodies capable of binding such ligands.

These and other objects of the invention will be apparent to the ordinary artisan upon consideration of the specification as a whole.

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#### SUMMARY OF THE INVENTION

The objects of the invention are achieved by providing an isolated mammalian megakaryocytopoietic proliferation and maturation promoting protein, denominated the "mpl ligand" (ML) or "thrombopoietin" (TPO), capable of stimulating proliferation, maturation and/or differentiation of megakaryocytes into the mature platelet-producing form.

This substantially homogeneous protein may be purified from a natural source by a method comprising; (1) contacting a source plasma containing the *mpl* ligand molecules to be purified with an immobilized receptor polypeptide, specifically *mpl* or a *mpl* fusion polypeptide immobilized on a support, under conditions whereby the *mpl* ligand molecules to be purified are selectively adsorbed onto the immobilized receptor polypeptide, (2) washing the immobilized receptor polypeptide and its support to remove non-adsorbed material, and (3) eluting the *mpl* ligand molecules from the immobilized receptor polypeptide to which they are adsorbed with an elution buffer. Preferably the natural source is mammalian plasma or urine containing the *mpl* ligand. Optionally the mammal is aplastic and the immobilized receptor is a *mpl*-IgG fusion.

Optionally, the prefered megakaryocytopoietic proliferation and maturation promoting protein is an isolated substantially homogeneous *mpl* ligand polypeptide made by synthetic or recombinant means.

The "mpl ligand" polypeptide or "TPO" of this invention preferably has at least 70% overall sequence identity with the amino acid sequence of the highly purified substantially homogeneous porcine mpl ligand polypeptide and at least 80% sequence identity with the "EPO-domain" of the porcine mpl ligand polypeptide. Optionally, the mpl ligand of this invention is mature human mpl ligand (hML), having the mature amino acid sequence provided in Fig. 1 (SEQ ID NO: 1), or a variant or posttranscriptionally modified form thereof or a protein having about 80% sequence identity with mature human mpl ligand. Optionally the mpl ligand variant is a fragment, especially an amino-terminus or "EPO-domain" fragment, of the mature human mpl ligand (hML). Preferably the amino terminus fragment retains substantially all of the human ML sequence between the first and forth cysteine residues but may contain substantial additions, deletions or substitutions outside that region. According to this embodiment, the fragment polypeptide may be represented by the formula:

#### X-hML(7-151)-Y

Where hML(7-151) represents the human TPO (hML) amino acid sequence from Cys<sup>7</sup> through Cys<sup>151</sup> inclusive; X represents the amino group of Cys<sup>7</sup> or one or more of the

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amino-terminus amino acid residue(s) of the mature hML or amino acid residue extensions thereto such as Met, Tyr or leader sequences containing, for example, proteolytic cleavage sites (e.g. Factor Xa or thrombin); and Y represents the carboxy terminal group of Cys<sup>151</sup> or one or more carboxy-terminus amino acid residue(s) of the mature hML or extensions thereto.

Optionally the *mpl* ligand polypeptide or fragment thereof may be fused to a heterologous polypeptide (chimera). A preferred heterologous polypeptide is a cytokine, colony stimulating factor or interleukin or fragment thereof, especially kitligand (KL), IL-1, IL-3, IL-6, IL-11, EPO, GM-CSF or LIF. An optional preferred heterologous polypeptide is an immunoglobin chain, especially human IgG1, IgG2, IgG3, IgG4, IgA, IgE, IgD, IgM or fragment thereof, especially comprising the constant domain of an IgG heavy chain.

Another aspect of this invention provides a composition comprising an isolated *mpl* agonist that is biologically active and is preferably capable of stimulating the incorporation of labeled nucleotides (e.g., <sup>3</sup>H-thymidine) into the DNA of IL-3 dependent Ba/F3 cells transfected with human *mpl*. Optionally the *mpl* agonist is biologically active *mpl* ligand and is preferably capable of stimulating the incorporation of <sup>35</sup>S into circulating platelets in a mouse platelet rebound assay. Suitable *mpl* agonist include hML153, hML(R153A, R154A), hML2, hML3, hML4, mML, mML2, mML3, pML, and pML2 or fragments thereof.

In another embodiment, this invention provides an isolated antibody capable of binding to the *mpl* ligand. The isolated antibody capable of binding to the *mpl* ligand may optionally be fused to a second polypeptide and the antibody or fusion thereof may be used to isolate and purify *mpl* ligand from a source as described above for immobilized *mpl*. In a further aspect of this embodiment, the invention provides a method for detecting the *mpl* ligand *in vitro* or *in vivo* comprising contacting the antibody with a sample, especially a serum sample, suspected of containing the ligand and detecting if binding has occurred.

In still further embodiments, the invention provides an isolated nucleic acid molecule, encoding the *mpl* ligand or fragments thereof, which nucleic acid molecule may optionally be labeled with a detectable moiety, and a nucleic acid molecule having a sequence that is complementary to, or hybridizes under moderate to highly stringent conditions with, a nucleic acid molecule having a sequence encoding a *mpl* ligand. Preferred nucleic acid molecules are those encoding human, porcine, and murine *mpl* ligand, and include RNA and DNA, both genomic and cDNA. In a further aspect of this embodiment, the nucleic acid molecule is DNA encoding the *mpl* ligand and further comprises a replicable vector in which the DNA is operably linked to control sequences

recognized by a host transformed with the vector. Optionally the DNA is cDNA having the sequence provided in Fig. 1 5'-3' (SEQ ID NO: 2), 3'-5' or a fragment thereof. This aspect further includes host cells, preferably CHO cells, transformed with the vector and a method of using the DNA to effect production of *mpl* ligand, preferably comprising expressing the cDNA encoding the *mpl* ligand in a culture of the transformed host cells and recovering the *mpl* ligand from the host cells or the host cell culture. The *mpl* ligand prepared in this manner is preferably human *mpl* ligand.

The invention further includes a method for treating a mammal having a hematopoietic disorder, especially thrombocytopenia, comprising administering a therapeutically effective amount of a *mpl* ligand to the mammal. Optionally the *mpl* ligand is administered in combination with a cytokine, especially a colony stimulating factor or interleukin. Preferred colony stimulating factors or interleukins include; kit-ligand (KL), LIF, G-CSF, GM-CSF, M-CSF, EPO, IL-1, IL-3, IL-6, and IL-11.

The invention further includes a process for isolating and purifying TPO (ML) from a TPO producing microorganism comprising:

(1) disrupting or lysing cells containing TPO.

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- (2) optionally seperating soluble material from insoluble material containing TPO,
  - (3) solublizing TPO in the insoluble material with a solublizing buffer,
  - (4) seperating solublized TPO from other soluble and insoluble material,
  - (5) refolding TPO in a redox buffer, and
  - (6) separating properly folded TPO from misfolded TPO.

The process provides for solubilizing the insoluble material containing TPO with a chaotropic agent where the chaotropic agent is selected from a salt of quanidine, sodium thiocyanate, or urea. The process further provides that solublized TPO is seperated from other soluble and insoluble material by one or more steps selected from centrafugation, gel filtration and reverse phase chromotography. The refolding step of the process provides for a redox buffer containing both an oxidizing and reducing agent. Generally, the oxidizing agent is oxygen or a compound containing at least one disulfide bond and the reducing agent is a compound containing at least one Preferably, the oxidizing agent is selected from oxidized free sulfhydryl. glutathione(GSSG) and cystine and the reducing agent is selected from reduced glutathione(GSH) and cysteine. Most preferably the oxidizing agent is oxidized glutathione(GSSG) and the reducing agent is reduced glutathione(GSH). It is also prefered that the molar ratio of the oxidizing agent is equal to or greater then that of the reducing agent. The redox buffer additionally contains a detergent, preferably selected from CHAPS and CHAPSO, present at a level of at least1%. The redox buffer additionally contains NaCl preferably at a concentration range of about 0.1-0.5M, and

glycerol preferably at a concentration greater than 15%. The pH of the redox buffer preferably ranges from about pH 7.5-pH 9.0. and the refolding step is conducted at 4 degrees for 12-48hr. The refolding step produces biologically active TPO in which a disulfide bond is formed between the Cys nearest the amino-terminus with the Cys nearest the carboxy-terminus of the EPO domain.

The invention further includes a process for purifying biologically active TPO from a microorganism comprising:

- (1) lysing at least the extracellular membrane of the microorganism,
  - (2) treating the lysate containing TPO with a chaotropic agent,
  - (3) refolding the TPO, and
- (4) separating impurities and misfolded TPO from properly folded TPO.

## BRIEF DESCRIPTION OF THE FIGURES

Fig. 1 shows the deduced amino acid sequence (SEQ ID NO: 1) of human *mpl* ligand (hML) cDNA and the coding nucleotide sequence (SEQ ID NO: 2). Nucleotides are numbered at the beginning of each line. The 5' and 3' untranslated regions are indicated in lower case letters. Amino acid residues are numbered above the sequence starting at Ser 1 of the mature *mpl* ligand (ML) protein sequence. The boundries of presumed exon 3 are indicated by the arrows and the potential N-glycosylation sites are boxed. Cysteine residues are indicated by a dot above the sequence. The underlined sequence corresponds to the N-terminal sequence determined from *mpl* ligand purified from porcine plasma.

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- Fig. 2 shows the procedure used for the *mpl* ligand  $^3$ H-thymidine incorporation assay. To determine the presence of *mpl* ligand from various sources, the *mpl* P Ba/F3 cells were starved of IL-3 for 24 hours in a humidified incubator at 37°C in 5% CO<sub>2</sub> and air. Following IL-3 starvation the cells were plated out in 96 well culture dishes with or without diluted samples and cultured for 24 hrs in a cell culture incubator. 20  $\mu$ l of serum free RPMI media containing 1  $\mu$ Ci of  $^3$ H-thymidine was added to each well for the last 6-8 hours. The cells were then harvested on 96 well filter plates and washed with water. The filters were then counted.
- Fig. 3 shows the effect of pronase, DTT and heat on the ability of APP to stimulate Ba/F3-mpl cell proliferation. For pronase digestion of APP, pronase (Boehringer Mannheim) or bovine serum albumin was coupled to Affi-gel10 (Biorad) and incubated individually with APP for 18hrs. at 37°C. Subsequently, the resins were

removed by centrifugation and supernatants assayed. APP was also heated to 80°C for 4 min. or made 100 µM DTT followed by dialysis against PBS.

Fig. 4 shows the elution of mpl ligand activity from Phenyl-Toyopearl, Blue-Sepharose and Ultralink-mpl columns. Fractions 4-8 from the mpl affinity column were the peak activity fractions eluted from the column.

Fig. 5 shows the SDS-PAGE of eluted Ultralink-*mpl* fractions. To 200 μl of each fraction 2-8, 1 ml of acetone containing 1mM HCl at -20°C was added. After 3hrs. at -20°C samples were centrifuged and resultant pellets were washed 2x with acetone at -20°C. The acetone pellets were subsequently dissolved in 30 μl of SDS-solubilization buffer, made 100 μM DTT and heated at 90°C for 5 min. The samples were then resolved on a 4-20% SDS-polyacrylamide gel and proteins were visualized by silver staining.

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Fig. 6 shows elution of *mpl* ligand activity from SDS-PAGE. Fraction 6 from the *mpl*-affinity column was resolved on a 4-20% SDS-polyacrylamide gel under non-reducing conditions. Following electrophoresis the gel was sliced into 12 equal regions and electroeluted as described in the examples. The electroeluted samples were dialyzed into PBS and assayed at a 1/20 dilution. The Mr standards used to calibrate the gel were Novex Mark 12 standards.

Fig. 7 shows the effect of *mpl* ligand depleted APP on human megakaryocytopoiesis. *mpl* ligand depleted APP was made by passing 1 ml over a 1 ml *mpl*-affinity column (700 μg *mpl*-lgG/ml NHS-superose, Pharmacia). Human peripheral stem cell cultures were made 10% APP or 10% *mpl* ligand depleted APP and cultured for 12 days. Megakaryocytopoiesis was quantitated as described in the examples.

Fig. 8 shows the effect of *mpl*-lgG on the stimulation of human megakaryocytopoiesis by APP. Human peripheral stem cell cultures were made 10% with APP and cultured for 12 days. At day 0, 2 and 4, *mpl*-lgG (0.5 μg) or ANP-R-lgG (0.5 μg) was added. After 12 days megakaryocytopoiesis was quantitated as described in the examples. The average of duplicate samples is graphed with the actual duplicate data in parenthesis.

Fig. 9 shows both strands of a 390 bp fragment of human genomic DNA encoding the mpl ligand. The deduced amino acid sequence of "exon 3" (SEQ ID NO: 3), the coding sequence (SEQ ID NO: 4), and its compliment (SEQ ID NO: 5) are shown.

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Fig. 10 shows deduced amino acid sequence of mature human *mpl* ligand (hML) (SEQ ID NO: 6) and mature human erythropoietin (hEPO) (SEQ ID NO: 7). The predicted amino acid sequence for the human *mpl* ligand is aligned with the human erythropoietin sequence. Identical amino acids are boxed and gaps introduced for optimal alignment are indicated by dashes. Potential N-glycosylation sites are underlined with a plain line for the hML and with a broken line for hEPO. The two cysteines important for erythropoietin activity are indicated by a large dot.

Fig. 11 shows deduced amino acid sequence of mature human *mpl* ligand isoforms hML (SEQ ID NO: 6), hML2 (SEQ ID NO: 8), hML3 (SEQ ID NO: 9), and hML4 (SEQ ID NO: 10). Identical amino acids are boxed and gaps introduced for optimal alignment are indicated by dashes.

Figs. 12A, 12B and 12C show the effect of human *mpl* ligand on Ba/F3-*mpl* cell proliferation (A), *in vitro* human megakaryocytopoiesis quantitated using a radiolabeled murine IgG monoclonal antibody specific to the megakaryocyte glycoprotein GPIIbIIIa (B), and murine thrombopoiesis measured in a platelet rebound assay (C).

Two hundred ninety-three cells were transfected by the CaPO4 method (Gorman, C in DNA Cloning: A New Approach 2:143-190 [1985]) with pRK5 vector alone, pRK5-hML or with pRK5-ML153 overnight (pRK5-ML153 was generated by introducing a stop codon after residue 153 of hML by PCR). Media was then conditioned for 36h and assayed for stimulation of cell proliferation of Ba/F3-mpl as described in Example 1 (A) or in vitro human megakaryocytopoiesis (B). Megakaryocytopoiesis was quantitated using a 1251 radiolabeled murine IgG monoclonal antibody (HP1-1D) to the megakaryocyte specific glycoprotein GPIIbIIIa as described (Grant et al., Blood 69:1334-1339 [1987]). The effect of partially purified recombinant ML (rML) on in vivo platelet production (C) was determined using the rebound thrombocytosis assay described by McDonald, T.P. Proc. Soc. Exp. Biol. Med. 144:1006-10012 (1973). Partially purified rML was prepared from 200ml of conditioned media containing the recombinant ML. The media was passed through a 2ml Blue-Separose column equilabrated in PBS and the column was washed with PBS and eluted with PBS containing 2M each of urea and NaCl. The active fraction was dialyzed into PBS and made 1mg/ml with endotoxin free BSA. The sample contained less than one unit of endotoxin /ml. Mice were injected with either 64,000, 32,000 or 16,000 units of rML or excipient alone. Each group consisted of six mice. The mean and standard deviation of each group is shown. p values were determined by a 2 tailed T-test comparing medians.

Fig. 13 compares the effect of human *mpl* ligand isoforms and variants in the Ba/F3-mpl cell proliferation assay. hML, mock, hML2, hML3, hML(R153A, R154A), and hML153 were assayed at various dilutions as described in **Example 1**.

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Figs. 14A, 14B and 14C show the deduced amino acid sequence (SEQ ID NO: 1) of human *mpl* ligand (hML) or human TPO (hTPO) and the human genomic DNA coding sequence (SEQ ID NO: 11). Nucleotides and amino acid residues are numbered at the beginning of each line.

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Fig. 15 shows a SDS-PAGE of purified 293-rhML332 and purified 293-rhML153.

Fig. 16 shows the nucleotide sequence: cDNA coding (SEQ ID NO: 12) and deduced amino acid sequence (SEQ ID NO: 13) of the open reading frame of a murine ML isoform. This mature murine mpl ligand isoform contains 331 amino acid residues, four fewer than the putative full length mML, and is therefore designated mML2. Nucleotides are numbered at the beginning of each line. Amino acid residues are numbered above the sequence starting with Ser 1. The potential N-glycosylation sites are underlined. Cysteine residues are indicated by a dot above the sequence.

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Fig. 17 shows the cDNA sequence (SEQ ID NO: 14) and predicted protein sequence (SEQ ID NO: 15) of this murine ML isoform (mML). Nucleotides are numbered at the beginning of each line. Amino acid residues are numbered above the sequence starting with Ser 1. This mature murine *mpl* ligand isoform contains 335 amino acid residues and is believed to be the full length *mpl* ligand, designated mML. The signal sequence is indicated with a dashed underline and the likely cleavage point is denoted with an arrow. The 5' and 3' untranslated regions are indicated with lower case letters. The two deleations found as a result of alternative splicing (mML2 and mML3) are underlined. The four cysteine residues are indicated by a dot. The seven potential N-glycosylation sites are boxed.

Fig. 18 compares the deduced amino acid sequence of the human ML isoform hML3 (SEQ ID NO: 9) and a murine ML isoform designated mML3 (SEQ ID NO: 16). The predicted amino acid sequence for the human *mpl* ligand is aligned with the murine *mpl* ligand sequence. Identical amino acids are boxed and gaps introduced for optimal alignment are indicated by dashes. Amino acids are numbered at the beginning of each line.

Fig. 19 compares the predicted amino acid sequences of mature ML isoforms from mouse-ML (SEQ ID NO: 17), porcine-ML (SEQ ID NO: 18) and human-ML (SEQ ID NO: 6). Amino acid sequences are aligned with gaps, indicated by dashes, introduced for optimal alingment. Amino acids are numbered at the beginning of each line with Identical residues boxed. Potential N-glycosylation sites are indicated by a shaded box and cysteine residues are designated with a dot. The conserved di-basic amino acid motif that presents a potential protease cleavage site is underlined. The four amino acid deletion found to occur in all three species (ML2) is outlined with a bold box.

Fig. 20 shows the cDNA sequence (SEQ ID NO: 19) and predicted mature protein sequence (SEQ ID NO: 18) of a porcine ML isoform (pML). This porcine mpl ligand isoform contains 332 amino acid residues and is believed to be the full length porcine mpl ligand, designated pML. Nucleotides are numbered at the beginning of each line. Amino acid residues are numbered above the sequence starting with Ser 1.

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Fig. 21 shows the cDNA sequence (SEQ ID NO: 20) and predicted mature protein sequence (SEQ ID NO: 21) of a porcine ML isoform (pML2). This porcine mpl ligand isoform contains 328 amino acid residues and is a four residues deletion form of the full length porcine mpl ligand, designated pML2. Nucleotides are numbered at the beginning of each line. Amino acid residues are numbered above the sequence starting with Ser 1.

Fig. 22 compares the deduced amino acid sequence of the full length porcine ML isoform pML (SEQ ID NO: 18) and a porcine ML isoform designated pML2 (SEQ ID NO: 21). The predicted amino acid sequence for the pML is aligned with pML2 sequence. Identical amino acids are boxed and gaps introduced for optimal alignment are indicated by dashes. Amino acids are numbered at the beginning of each line.

Fig. 23 shows the pertinent features of plasmid pSVI5.ID.LL.MLORF ("full length" or TPO332) used to transfect host CHO-DP12 cells for production of CHO-rhTPO332.

Fig. 24 shows the pertinent features of plasmid pSVI5.ID.LL.MLEPO-D ("truncated" or TPO153) used to transfect host CHO-DP12 cells for production of CHO-rhTPO153.

Figs. 25A, 25B, and 25C show the effect of *E. coli*-rhTPO(Met<sup>-1</sup>, 153) on platelets (A), red blood cells (B) and (C) white blood cells in normal mice. Two groups of 6 female C57 B6 mice were injected daily with either PBS buffer or 0.3μg *E. coli*-rhTPO(Met<sup>-1</sup>, 153) (100μl sc.). On day 0 and on days 3-7 40μl of blood was

taken from the orbital sinus. This blood was immediately diluted in 10 ml of commercial diluant and complete blood counts were obtained on a Serrono Baker Hematology Analyzer 9018. The data are presented as means  $\pm$  Standard error of the mean.

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Figs. 26A, 26B and 26C show the effect of *E. coli*-rhTPO(Met<sup>-1</sup>, 153) on platelets (A), red blood cells (B) and (C) white blood cells in sublethally irradiated mice. Two groups of 10 female C57 B6 mice were sublethally irradiated with 750 cGy of gamma radiation from a <sup>137</sup>Cs source and injected daily with either PBS buffer or 3.0μg *E. coli*-rhTPO(Met<sup>-1</sup>, 153) (100μl sc.). On day 0 and at subsequent intermediate time points 40μl of blood was taken from the orbital sinus. This blood was immediately diluted in 10 ml of commercial diluant and complete blood counts were obtained on a Serrono Baker Hematology Analyzer 9018. The data are presented as means ± Standard error of the mean.

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Figs. 27A, 27B and 27C show the effect of CHO-rhTPO332 on (A) platelets (thrombocytes), (B) red blood cells (erythrocytes) and (C) white blood cells (leukocytes) in normal mice. Two groups of 6 female C57 B6 mice were injected daily with either PBS buffer or  $0.3\mu g$  CHO-rhTPO332 ( $100\mu l$  sc.). On day 0 and on days 3-7  $40\mu l$  of blood was taken from the orbital sinus. This blood was immediately diluted in 10 ml of commercial diluant and complete blood counts were obtained on a Serrono Baker Hematology Analyzer 9018. The data are presented as means  $\pm$  Standard error of the mean.

Fig. 28 shows dose response curves for various forms of rhTPO obtained from various cell lines. Dose response curves were constructed to rhTPO from the following cell lines: hTPO332 from CHO (full length from Chinese hamster ovary cells); hTPO<sub>Met</sub>-1 153 (E. coli-derived truncated form with an N-terminal methionine); hTPO332 (full length TPO from human 293 cells); Met-less 155 E-Coli (the truncated form [rhTPO155] without the terminal methionine from E. coli). Groups of 6 female C57B6 mice were injected daily for 7 days with rhTPO depending upon group. Each day 40μl of blood was taken from the orbital sinus for complete blood counts. The data presented above are the maximal effects seen with the various treatments and with the exception of (met 153 E-Coli) this occurred on day 7 of treatment. In the aforementioned "met 153 E-Coli" group the maximal effect was seen on day 5. The data are presented as means ± Standard error of the mean.

Fig. 29 shows dose response curves comparing the activity of full length and "cliped" forms of rhTPO produced in CHO cells with the truncated form from *E. coli*. Groups of 6 female C57B6 mice were injected daily with 0.3μg rhTPO of various types. On days 2-7 40μl of blood was taken from the orbital sinus for complete blood counts. Treatment groups were TPO<sub>153</sub> the truncated form of TPO from *E. coli*; TPO<sub>332</sub> (Mix fraction) Full length TPO containing approximately 80-90% full length and 10-20% clipped forms; TPO<sub>332</sub>(30K fraction) = purified clipped fraction from the original "mix" preparation; TPO<sub>332</sub>(70K fraction) = purified full length TPO fraction from the original "mix" preparation. The data are presented as means ± Standard error of the mean.

Fig. 30 is a cartoon showing the KIRA ELISA assay for measuring TPO. The figure shows the MPL/Rse.gD chimera and relavant parts of the parent receptors as well as the final construct (right portion of the figure) and a flow diagram (left portion of the figure) showing relavant steps of the assay.

Fig. 31 is a flow chart for the KIRA ELISA assay showing each step in the procedure.

Figs. 32A-32L provide the nucleotide sequence (SEQ ID NO: 22) of the pSVI17.ID.LL expression vector used for expression of Rse.gD in Example 17.

Fig. 33 is a schematic representation of the preparation of plasmid pMP1.

Fig. 34 is a schematic representation of the preperation of plasmid pMP21.

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Fig. 35 is a schematic representation of the preparation of plasmid pMP151.

Fig. 36 is a schematic representation of the preperation of plasmid pMP202.

30 Fig. 37 is a schematic representation of the preperation of plasmid pMP172.

Fig. 38 is a schematic representation of the preperation of plasmid pMP210.

Fig. 39 is a table of the five best expressing TPO clones from the pMP210 plasmid bank (SEQ ID NOS: 23, 24, 25, 26, 27 and 28).

Fig. 40 is a schematic representation of the preperation of plasmid pMP41.

Fig. 41 is a schematic representation of the preperation of plasmid pMP57.

Fig. 42 is a schematic representation of the preperation of plasmid pMP251.

#### DETAILED DESCRIPTION OF THE INVENTION

#### I. Definitions

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In general, the following words or phrases have the indicated definition when used in the description, examples, and claims.

"Chaotropic agent" refers to a compound which, in aqueous solution and in suitable concentrations, can cause a change in the spatial configuration or conformation of a protein by at least partially disrupting the forces responsible for maintaining the normal secondary and tertiary structure of the protein. Such compounds include, for example, urea, guanidine HCI, and sodium thiocyanate. High concentrations, usually 4-9M, of these compounds are normally required to exert the conformational effect on proteins.

"Cytokine" is a generic term for proteins released by one cell population which act on another cell as intercellular mediators. Examples of such cytokines are lymphokines, monokines, and traditional polypeptide hormones. Included among the cytokines are growth hormone, insulin-like growth factors, human growth hormone, N-methionyl human growth hormone, bovine growth hormone, parathyroid hormone, thyroxine, insulin, proinsulin, relaxin, prorelaxin, glycoprotein hormones such as follicle stimulating hormone (FSH), thyroid stimulating hormone (TSH), and leutinizing hormone (LH), hematopoietic growth factor, hepatic growth factor, fibroblast growth factor, prolactin, placental lactogen, tumor necrosis factor- $\alpha$  $(TNF-\alpha \ and \ TNF-\beta)$  mullerian-inhibiting substance, mouse gonadotropin-associated peptide, inhibin, activin, vascular endothelial growth factor, integrin, nerve growth factors such as NGF-β, platelet-growth factor, transforming growth factors (TGFs) such as TGF-α and TGF-β, insulin-like growth factor-I and -II, erythropoletin (EPO), osteoinductive factors, interferons such as interferon- $\alpha$ , - $\beta$ , and - $\gamma$ , colony stimulating factors (CSFs) such as macrophage-CSF (M-CSF), granulocytemacrophage-CSF (GM-CSF), and granulocyte-CSF (G-CSF), interleukins (IL's) such as IL-1, IL-1 $\alpha$ , IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-11, IL-12 and other polypeptide factors including LIF, SCF, and kit-ligand. As used herein the foregoing terms are meant to include proteins from natural sources or from recombinant cell culture. Similarly, the terms are intended to include biologically active equivalents; e.g., differing in amino acid sequence by one or more amino acids or in type or extent of glycosylation.

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"mpl ligand", "mpl ligand polypeptide", "ML", "thrombopoletin" or "TPO" are used interchangeably herein and comprise any polypeptide that possesses the property of binding to mpl, a member of the cytokine receptor superfamily, and having a biological property of the ML as defined below. An exemplary biological property is the ability to stimulate the incorporation of labeled nucleotides (e.g., <sup>3</sup>H-thymidine) into the DNA of IL-3 dependent Ba/F3 cells transfected with human mpl P. Another exemplary biological property is the ability to stimulate the incorporation of <sup>35</sup>S into circulating platelets in a mouse platelet rebound assay. This definition encompasses the polypeptide isolated from a mpl ligand source such as aplastic porcine plasma described herein or from another source, such as another animal species, including humans or prepared by recombinant or synthetic methods and includes variant forms including functional derivatives, fragments, alleles, isoforms and analogues thereof.

A "mpl ligand fragment" or "TPO fragment" is a portion of a naturally occurring mature full length mpl ligand or TPO sequence having one or more amino acid residues or carbohydrate units deleted. The deleted amino acid residue(s) may occur anywhere in the peptide including at either the N-terminal or C-terminal end or internally. The fragment will share at least one biological property in common with mpl ligand. Mpl ligand fragments typically will have a consecutive sequence of at least 10, 15, 20, 25, 30, or 40 amino acid residues that are identical to the sequences of the mpl ligand isolated from a mammal including the ligand isolated from aplastic porcine plasma or the human or murine ligand, especially the EPO-domain thereof. Representative examples of N-terminal fragments are hML153 or TPO(Met<sup>-1</sup>1-153).

"Mpl ligand variants" or "mpl ligand sequence variants" as defined herein means a biologically active mpl ligand as defined below having less than 100% sequence identity with the mpl ligand isolated from recombinant cell culture or aplastic porcine plasma or the human ligand having the deduced sequence described in Fig. 1 (SEQ ID NO: 1). Ordinarily, a biologically active mpl ligand variant will have an amino acid sequence having at least about 70% amino acid sequence identity with the mpl ligand isolated from aplastic porcine plasma or the mature murine or human ligand or fragments thereof (see Fig. 1 [SEQ ID NO: 1]), preferably at least about 75%, more preferably at least about 80%, still more preferably at least about 95%.

A "chimeric *mpl* ligand" is a polypeptide comprising full length *mpl* ligand or one or more fragments thereof fused or bonded to a second heterologous polypeptide or one or more fragments thereof. The chimera will share at least one biological property in common with *mpl* ligand. The second polypeptide will typically be a cytokine, immunoglobin or fragment thereof.

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"Isolated *mpl* ligand", "highly purified *mpl* ligand" and "substantially homogeneous *mpl* ligand" are used interchangeably and mean a *mpl* ligand that has been purified from a *mpl* ligand source or has been prepared by recombinant or synthetic methods and is sufficiently free of other peptides or proteins (1) to obtain at least 15 and preferably 20 amino acid residues of the N-terminal or of an internal amino acid sequence by using a spinning cup sequenator or the best commercially available amino acid sequenator marketed or as modified by published methods as of the filing date of this application, or (2) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Homogeneity here means less than about 5% contamination with other source proteins.

"Biological property" when used in conjunction with either the "mpl ligand" or "Isolated mpl ligand" means having thrombopoletic activity or having an in vivo effector or antigenic function or activity that is directly or indirectly caused or performed by a mpl ligand (whether in its native or denatured conformation) or a fragment thereof. Effector functions include mpl binding and any carrier binding activity, agonism or antagonism of mpl, especially transduction of a proliferative signal including replication, DNA regulatory function, modulation of the biological activity of other cytokines, receptor (especially cytokine) activation, deactivation, up- or down regulation, cell growth or differentiation and the like. An antigenic function means possession of an epitope or antigenic site that is capable of crossreacting with antibodies raised against the native mpl ligand. The principal antigenic function of a mpl ligand polypeptide is that it binds with an affinity of at least about 106 l/mole to an antibody raised against the mpl ligand isolated from aplastic porcine plasma. Ordinarily, the polypeptide binds with an affinity of at least about 107 Most preferably, the antigenically active mpl ligand polypeptide is a polypeptide that binds to an antibody raised against the mpl ligand having one of the above described effector functions. The antibodies used to define "biologically activity" are rabbit polyclonal antibodies raised by formulating the mpl ligand isolated from recombinant cell culture or aplastic porcine plasma in Freund's complete adjuvant, subcutaneously injecting the formulation, and boosting the immune response by intraperitoneal injection of the formulation until the titer of mpl ligand antibody plateaus.

"Isolated *mpl* ligand" means a *mpl* ligand or polypeptide that exhibits thrombopoietic activity or shares an effector function of the *mpl* ligand isolated from aplastic porcine plasma or expressed in recombinant cell culture described herein. A principal known effector function of the *mpl* ligand or polypeptide herein is binding to *mpl* and stimulating the incorporation of labeled nucleotides (<sup>3</sup>H-thymidine) into the DNA of

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IL-3 dependent Ba/F3 cells transfected with human *mpl* P. Another known effector function of the *mpl* ligand or polypeptide herein is the ability to stimulate the incorporation of <sup>35</sup>S into circulating platelets in a mouse platelet rebound assay. Yet another known effector function of *mpl* ligand is the ability to stimulate *in vitro* human megakaryocytopolesis that may be quantitated by using a radio labeled monoclonal antibody specific to the megakaryocyte glycoprotein GPII<sub>b</sub>III<sub>a</sub>.

"Percent amino acid sequence identity" with respect to the *mpl* ligand sequence is defined herein as the percentage of amino acid residues in the candidate sequence that are identical with the residues in the *mpl* ligand sequence isolated from aplastic porcine plasma or the murine or human ligand having the deduced amino acid sequence described in Fig. 1 (SEQ ID NO: 1), after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. None of N-terminal, C-terminal, or internal extensions, deletions, or insertions into the *mpl* ligand sequence shall be construed as affecting sequence identity or homology. Thus exemplary biologically active *mpl* ligand polypeptides considered to have identical sequences include; prepro-*mpl* ligand, pro-*mpl* ligand, and mature *mpl* ligand.

"Mpl ligand microsequencing" may be accomplished by any appropriate standard procedure provided the procedure is sensitive enough. In one such method, highly purified polypeptide obtained from SDS gels or from a final HPLC step are sequenced directly by automated Edman (phenyl isothiocyanate) degradation using a model 470A Applied Biosystems gas phase sequencer equipped with a 120A phenylthiohydantion (PTH) amino acid analyzer. Additionally, mpl ligand fragments prepared by chemical (e.g., CNBr, hydroxylamine, 2-nitro-5-thiocyanobenzoate) or enzymatic (e.g., trypsin, clostripain, staphylococcal protease) digestion followed by fragment purification (e.g., HPLC) may be similarly sequenced. PTH amino acids are analyzed using the ChromPerfect data system (Justice Innovations, Palo Alto, CA). Sequence interpretation is performed on a VAX 11/785 Digital Equipment Co. computer as described by Henzel et al., J. Chromatography, 404:41-52 [1987]. Optionally, aliquots of HPLC fractions may be electrophoresed on 5-20% SDS-PAGE, electrotransferred to a PVDF membrane (ProBlott, AIB, Foster City, CA) and stained with Coomassie Brilliant Blue (Matsurdiara, J. Biol. Chem., 262:10035-10038 [1987]. A specific protein identified by the stain is excised from the blot and Nterminal sequencing is carried out with the gas phase sequenator described above. For internal protein sequences, HPLC fractions are dried under vacuum (SpeedVac), resuspended in appropriate buffers, and digested with cyanogen bromide, the Lysspecific enzyme Lys-C (Wako Chemicals, Richmond, VA), or Asp-N (Boehringer Mannheim, Indianapolis, IN). After digestion, the resultant peptides are sequenced as a

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mixture or after HPLC resolution on a C4 column developed with a propanol gradient in 0.1% TFA prior to gas phase sequencing.

"Thrombocytopenia" is defined as a platelet count below 150  $\times$  10 $^9$  per liter of blood.

"Thrombopoietic activity" is defined as biological activity that consists of accelerating the proliferation, differentiation and/or maturation of megakaryocytes or megakaryocyte precursors into the platelet producing form of these cells. This activity may be measured in various assays including an *in vivo* mouse platelet rebound synthesis assay, induction of platelet cell surface antigen assay as measured by an anti-platelet immunoassay (anti-GPIIbIIIa) for a human leukemia megakaryoblastic cell line (CMK), and induction of polyploidization in a megakaryoblastic cell line (DAMI).

"Thrombopoietin" (TPO) is defined as a compound having thrombopoietic activity or being capable of increasing serum platelet counts in a mammal. TPO is preferably capable of increasing endogenous platelet counts by at least 10%, more preferably by 50%, and most preferably capable of elevating platelet counts in a human to greater that 150X10<sup>9</sup> per liter of blood.

"Isolated *mpl* ligand nucleic acid" is RNA or DNA containing greater than 16 and preferably 20 or more sequential nucleotide bases that encode biologically active *mpl* ligand or a fragment thereof, is complementary to the RNA or DNA, or hybridizes to the RNA or DNA and remains stably bound under moderate to stringent conditions. This RNA or DNA is free from at least one contaminating source nucleic acid with which it is normally associated in the natural source and preferably substantially free of any other mammalian RNA or DNA. The phrase "free from at least one contaminating source nucleic acid with which it is normally associated" includes the case where the nucleic acid is present in the source or natural cell but is in a different chromosomal location or is otherwise flanked by nucleic acid sequences not normally found in the source cell. An example of isolated *mpl* ligand nucleic acid is RNA or DNA that encodes a biologically active *mpl* ligand sharing at least 75% sequence identity, more preferably at least 85%, even more preferably 90%, and most preferably 95% sequence identity with the human, murine or porcine *mpl* ligand.

"Control sequences" when referring to expression means DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, a ribosome binding site, and possibly, other as yet poorly understood sequences. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

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"Operably linked" when referring to nucleic acids means that the nucleic acids are placed in a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accord with conventional practice.

"Exogenous" when referring to an element means a nucleic acid sequence that is foreign to the cell, or homologous to the cell but in a position within the host cell nucleic acid in which the element is ordinarily not found.

"Cell," "cell line," and "cell culture" are used interchangeably herein and such designations include all progeny of a cell or cell line. Thus, for example, terms like "transformants" and "transformed cells" include the primary subject cell and cultures derived therefrom without regard for the number of transfers. It is also understood that all progeny may not be precisely identical in DNA content, due to deliberate or inadvertent mutations. Mutant progeny that have the same function or biological activity as screened for in the originally transformed cell are included. Where distinct designations are intended, it will be clear from the context.

"Plasmids" are autonomously replicating circular DNA molecules possessing independent origins of replication and are designated herein by a lower case "p" preceded and/or followed by capital letters and/or numbers. The starting plasmids herein are either commercially available, publicly available on an unrestricted basis, or can be constructed from such available plasmids in accordance with published procedures. In addition, other equivalent plasmids are known in the art and will be apparent to the ordinary artisan.

"Restriction enzyme digestion" when referring to DNA means catalytic cleavage of internal phosphodiester bonds of DNA with an enzyme that acts only at certain locations or sites in the DNA sequence. Such enzymes are called "restriction endonucleases". Each restriction endonuclease recognizes a specific DNA sequence called a "restriction site" that exhibits two-fold symmetry. The various restriction enzymes used herein are commercially available and their reaction conditions, cofactors, and other requirements as established by the enzyme suppliers are used. Restriction enzymes commonly are designated by abbreviations composed of a capital

letter followed by other letters representing the microorganism from which each restriction enzyme originally was obtained and then a number designating the particular enzyme. In general, about 1 µg of plasmid or DNA fragment is used with about 1-2 units of enzyme in about 20 µl of buffer solution. Appropriate buffers and substrate amounts for particular restriction enzymes are specified by the manufacturer. Incubation of about 1 hour at 37°C is ordinarily used, but may vary in accordance with the supplier's instructions. After incubation, protein or polypeptide is removed by extraction with phenol and chloroform, and the digested nucleic acid is recovered from the aqueous fraction by precipitation with ethanol. Digestion with a restriction enzyme may be followed with bacterial alkaline phosphatase hydrolysis of the terminal 5' phosphates to prevent the two restriction-cleaved ends of a DNA fragment from "circularizing" or forming a closed loop that would impede insertion of another DNA fragment at the restriction site. Unless otherwise stated, digestion of plasmids is not followed by 5' terminal dephosphorylation. Procedures and reagents for dephosphorylation are conventional as described in sections 1.56-1.61 of Sambrook et al., Molecular Cloning: A Laboratory Manual [New York: Cold Spring Harbor Laboratory Press, 1989].

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"Recovery" or "isolation" of a given fragment of DNA from a restriction digest means separation of the digest on polyacrylamide or agarose gel by electrophoresis, identification of the fragment of interest by comparison of its mobility versus that of marker DNA fragments of known molecular weight, removal of the gel section containing the desired fragment, and separation of the gel from DNA. This procedure is known generally. For example, see Lawn et al., Nucleic Acids Res., 9:6103-6114 [1981], and Goeddel et al., Nucleic Acids Res., 8:4057 [1980].

"Southern analysis" or "Southern blotting" is a method by which the presence of DNA sequences in a restriction endonuclease digest of DNA or DNA-containing composition is confirmed by hybridization to a known, labeled oligonucleotide or DNA fragment. Southern analysis typically involves electrophoretic separation of DNA digests on agarose gels, denaturation of the DNA after electrophoretic separation, and transfer of the DNA to nitrocellulose, nylon, or another suitable membrane support for analysis with a radiolabeled, biotinylated, or enzyme-labeled probe as described in sections 9.37-9.52 of Sambrook *et al.*, *supra*.

"Northern analysis" or "Northern blotting" is a method used to identify RNA sequences that hybridize to a known probe such as an oligonucleotide, DNA fragment, cDNA or fragment thereof, or RNA fragment. The probe is labeled with a radioisotope such as <sup>32</sup>P, or by biotinylation, or with an enzyme. The RNA to be analyzed is usually electrophoretically separated on an agarose or polyacrylamide gel, transferred to nitrocellulose, nylon, or other suitable membrane, and hybridized with the probe,

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using standard techniques well known in the art such as those described in sections 7.39-7.52 of Sambrook et al., supra.

"Ligation" is the process of forming phosphodiester bonds between two nucleic acid fragments. For ligation of the two fragments, the ends of the fragments must be compatible with each other. In some cases, the ends will be directly compatible after endonuclease digestion. However, it may be necessary first to convert the staggered ends commonly produced after endonuclease digestion to blunt ends to make them compatible for ligation. For blunting the ends, the DNA is treated in a suitable buffer for at least 15 minutes at 15°C with about 10 units of the Klenow fragment of DNA polymerase I or T4 DNA polymerase in the presence of the four deoxyribonucleotide triphosphates. The DNA is then purified by phenol-chloroform extraction and ethanol precipitation. The DNA fragments that are to be ligated together are put in solution in about equimolar amounts. The solution will also contain ATP, ligase buffer, and a ligase such as T4 DNA ligase at about 10 units per 0.5 μg of DNA. If the DNA is to be ligated into a vector, the vector is first linearized by digestion with the appropriate restriction endonuclease(s). The linearized fragment is then treated with bacterial alkaline phosphatase or calf intestinal phosphatase to prevent self-ligation during the ligation step.

"Preparation" of DNA from cells means isolating the plasmid DNA from a culture of the host cells. Commonly used methods for DNA preparation are the large-and small-scale plasmid preparations described in sections 1.25-1.33 of Sambrook et al., supra. After preparation of the DNA, it can be purified by methods well known in the art such as that described in section 1.40 of Sambrook et al., supra.

"Oligonucleotides" are short-length, single- or double-stranded polydeoxynucleotides that are chemically synthesized by known methods (such as phosphotriester, phosphite, or phosphoramidite chemistry, using solid-phase techniques such as described in EP 266,032 published 4 May 1988, or via deoxynucleoside H-phosphonate intermediates as described by Froehler *et al.*, *Nucl. Acids Res.*, 14:5399-5407 [1986]). Further methods include the polymerase chain reaction defined below and other autoprimer methods and oligonucleotide syntheses on solid supports. All of these methods are described in Engels *et al.*, *Agnew. Chem. Int. Ed. Engl.*, 28:716-734 (1989). These methods are used if the entire nucleic acid sequence of the gene is known, or the sequence of the nucleic acid complementary to the coding strand is available. Alternatively, if the target amino acid sequence is known, one may infer potential nucleic acid sequences using known and preferred coding residues for each amino acid residue. The oligonucleotides are then purified on polyacrylamide gels.

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"Polymerase chain reaction" or "PCR" refers to a procedure or technique in which minute amounts of a specific piece of nucleic acid, RNA and/or DNA, are amplified as described in U.S. Patent No. 4,683,195 issued 28 July 1987. Generally, sequence information from the ends of the region of interest or beyond needs to be avallable, such that oligonucleotide primers can be designed; these primers will be identical or similar in sequence to opposite strands of the template to be amplified. The 5' terminal nucleotides of the two primers may coincide with the ends of the amplified material. PCR can be used to amplify specific RNA sequences, specific DNA sequences from total genomic DNA, and cDNA transcribed from total cellular RNA, bacteriophage or plasmid sequences, etc. See generally Mullis et al., Cold Spring Harbor Symp. Quant. Biol., 51:263 [1987]; Erlich, ed., PCR Technology, (Stockton Press, NY, 1989). As used herein, PCR is considered to be one, but not the only, example of a nucleic acid polymerase reaction method for amplifying a nucleic acid test sample comprising the use of a known nucleic acid as a primer and a nucleic acid polymerase to amplify or generate a specific piece of nucleic acid.

"Stringent conditions" are those that (1) employ low ionic strength and high temperature for washing, for example, 0.015 M NaCl/0.0015 M sodium citrate/0.1% NaDodSO<sub>4</sub> (SDS) at 50°C, or (2) employ during hybridization a denaturing agent such as formamide, for example, 50% (vol/vol) formamide with 0.1% bovine serum albumin/0.1% Ficoll/0.1% polyvinylpyrrolidone/50 mM sodium phosphate buffer at pH 6.5 with 750 mM NaCl, 75 mM sodium citrate at 42°C. Another example is use of 50% formamide, 5 x SSC (0.75 M NaCl, 0.075 M sodium citrate), 50 mM sodium phosphate (pH 6.8), 0.1% sodium pyrophosphate, 5 x Denhardt's solution, sonicated salmon sperm DNA (50  $\mu$ g/ml), 0.1% SDS, and 10% dextran sulfate at 42°C, with washes at 42°C in 0.2 x SSC and 0.1% SDS.

"Moderately stringent conditions" are described in Sambrook *et al.*, *supra*, and include the use of a washing solution and hybridization conditions (*e.g.*, temperature, ionic strength, and %SDS) less stringent than described above. An example of moderately stringent conditions are conditions such as overnight incubation at 37°C in a solution comprising: 20% formamide, 5 X SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5 X Denhardt's solution, 10% dextran sulfate, and 20 µl/ml denatured sheared salmon sperm DNA, followed by washing the filters in 1 X SSC at about 37-50°C. The skilled artisan will recognize how to adjust the temperature, ionic strength *etc.* as necessary to accommodate factors such as probe length and the like.

"Antibodies" (Abs) and "immunoglobulins" (Igs) are glycoproteins having the same structural characteristics. While antibodies exhibit binding specificity to a specific antigen, immunoglobulins include both antibodies and other antibody-like

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molecules which lack antigen specificity. Polypeptides of the latter kind are, for example, produced at low levels by the lymph system and at increased levels by myelomas.

"Native antibodies and immunoglobulins" are usually heterotetrameric glycoproteins of about 150,000 dattons, composed of two identical light (L) chains and two identical heavy (H) chains. Each light chain is linked to a heavy chain by one covalent disulfide bond, while the number of disulfide linkages varies between the heavy chains of different immunoglobulin isotypes. Each heavy and light chain also has regularly spaced intrachain disulfide bridges. Each heavy chain has at one end a variable domain (V<sub>H</sub>) followed by a number of constant domains. Each light chain has a variable domain at one and (V<sub>L</sub>) and a constant domain at its other end; the constant domain of the light chain is aligned with the first constant domain of the heavy chain, and the light chain variable domain is aligned with the variable domain of the heavy chain. Particular amino acid residues are believed to form an interface between the light and heavy chain variable domains (Clothia *et al.*, *J.' Mol. Biol.*, 186:651-663 [1985]; Novotny and Haber, *Proc. Natl. Acad. Sci. USA*, 82:4592-4596 [1985]).

The term "variable" refers to the fact that certain portions of the variable domains differ extensively in sequence among antibodies and are used in the binding and specificity of each particular antibody for its particular antigen. However, the variability is not evenly distributed through the variable domains of antibodies. It is concentrated in three segments called complementarity determining regions (CDRs) or hypervariable regions both in the light chain and the heavy chain variable domains. The more highly conserved portions of variable domains are called the framework (FR). The variable domains of native heavy and light chains each comprise four FR regions, largely adopting a β-sheet configuration, connected by three CDRs, which form loops connecting, and in some cases forming part of, the  $\beta$ -sheet structure. The CDRs in each chain are held together in close proximity by the FR regions and, with the CDRs from the other chain, contribute to the formation of the antigen binding site of antibodies (see Kabat et al., Sequences of Proteins of Immunological Interest, National Institute of Health, Bethesda, MD [1987]). The constant domains are not involved directly in binding an antibody to an antigen, but exhibit various effector functions, such as participation of the antibody in antibody-dependent cellular toxicity.

Papain digestion of antibodies produces two identical antigen binding fragments, called "Fab" fragments, each with a single antigen binding site, and a residual "Fc" fragment, whose name reflects its ability to crystallize readily. Pepsin treatment yields an F(ab')<sub>2</sub> fragment that has two antigen combining sites and is still capable of cross-linking antigen.

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"Fv" is the minimum antibody fragment which contains a complete antigen recognition and binding site. This region consists of a dimer of one heavy and one light chain variable domain in tight, non-covalent association. It is in this configuration that the three CDRs of each variable domain interact to define an antigen binding site on the surface of the V<sub>H</sub>-V<sub>L</sub> dimer. Collectively, the six CDRs confer antigen binding specificity to the antibody. However, even a single variable domain (or half of an Fv comprising only three CDRs specific for an antigen) has the ability to recognize and bind antigen, although at a lower affinity than the entire binding site.

The Fab fragment also contains the constant domain of the light chain and the first constant domain (CH1) of the heavy chain. Fab' fragments differ from Fab fragments by the addition of a few residues at the carboxy terminus of the heavy chain CH1 domain including one or more cysteines from the antibody hinge region. Fab'-SH is the designation herein for Fab' in which the cysteine residue(s) of the constant domains bear a free thiol group. F(ab')<sub>2</sub> antibody fragments originally were produced as pairs of Fab' fragments which have hinge cysteines between them. Other, chemical couplings of antibody fragments are also known.

The "light chains" of antibodies (immunoglobulins) from any vertebrate species can be assigned to one of two clearly distinct types, called kappa and lambda  $(\lambda)$ , based on the amino acid sequences of their constant domains.

Depending on the amino acid sequence of the constant domain of their heavy chains, immunoglobulins can be assigned to different classes. There are five major classes of immunoglobulins: IgA, IgD, IgE, IgG and IgM, and several of these may be further divided into subclasses (isotypes), e.g., IgG-1, IgG-2, IgG-3, and IgG-4; IgA-1 and IgA-2. The heavy chain constant domains that correspond to the different classes of immunoglobulins are called  $\alpha$ , delta, epsilon,  $\gamma$ , and  $\mu$ , respectively. The subunit structures and three-dimensional configurations of different classes of immunoglobulins are well known.

The term "antibody" is used in the broadest sense and specifically covers single monoclonal antibodies (including agonist and antagonist antibodies), antibody compositions with polyepitopic specificity, as well as antibody fragments (e.g., Fab, F(ab')<sub>2</sub>, and Fv), so long as they exhibit the desired biological activity.

The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, *i.e.*, the individual antibodies comprising the population are identical except for possible naturally occurring mutations that may be present in minor amounts. Monoclonal antibodies are highly specific, being directed against a single antigenic site. Furthermore, in contrast to conventional (polyclonal) antibody preparations which typically include different antibodies directed against different determinants (epitopes), each

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monoclonal antibody is directed against a single determinant on the antigen. In addition to their specificity, the monoclonal antibodies are advantageous in that they are synthesized by the hybridoma culture, uncontaminated by other immunoglobulins. The modifier "monoclonal" indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies, and is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies to be used in accordance with the present invention may be made by the hybridoma method first described by Kohler & Milstein, *Nature*, 256:495 (1975), or may be made by recombinant DNA methods (see, *e.g.*, U.S. Patent No. 4,816,567 [Cabilly *et al.*]).

The monoclonal antibodies herein specifically include "chimeric" antibodies (immunoglobulins) in which a portion of the heavy and/or light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired biological activity (U.S. Patent No. 4,816,567 (Cabilly et al.); and Morrison et al., Proc. Natl. Acad. Sci. USA, 81:6851-6855 [1984]).

"Humanized" forms of non-human (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, humanized antibody may comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. These modifications are made to further refine and optimize antibody performance. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. For further details see: Jones

et al., Nature, 321:522-525 [1986]; Reichmann et al., Nature, 332:323-329 [1988]; and Presta, Curr. Op. Struct. Biol., 2:593-596 [1992]).

"Non-immunogenic in a human" means that upon contacting the polypeptide in a pharmaceutically acceptable carrier and in a therapeutically effective amount with the appropriate tissue of a human, no state of sensitivity or resistance to the polypeptide is demonstratable upon the second administration of the polypeptide after an appropriate latent period (e.g., 8 to 14 days).

### 11. Preferred Embodiments of the invention

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Preferred polypeptides of this invention are substantially homogeneous polypeptide(s), referred to as mpl ligand(s) or thrombopoletin (TPO), that possesse the property of binding to mpl, a member of the receptor cytokine superfamily, and having the biological property of stimulating the incorporation of labeled nucleotides (3H-thymidine) into the DNA of IL-3 dependent Ba/F3 cells transfected with human mpi P. More preferred mpi ligand(s) are isolated mammalian protein(s) having hematopoletic, especially megakaryocytopoletic or thrombocytopoletic activity namely, being capable of stimulating proliferation, maturation and/or differentiation of immature megakaryocytes or their predecessors into the mature platelet-producing Most preferred polypeptides of this invention are human mpl ligand(s) including fragments thereof having hematopoietic, megakaryocytopoietic or thrombopoietic activity. Optionally these human mpl ligand(s) lack glycosylation. Other prefered human mpl ligands are the "EPO-domain" of hML refered to as hML153 or hTPO153, a truncated form of hML refered to as hML245 or hTPO245 and the mature full length polypeptide having the amino acid sequence shown in Fig. 1 (SEQ ID NO: 1), refered to as hML, hML332 or hTPO332 and the biolocically active substitutional variant hML(R153A, R154A).

Optional preferred polypeptides of this invention are biologically or immunologically active *mpl* ligands variants selected from hML2, hML3, hML4, mML, mML2, mML3, pML and pML2.

Optional preferred polypeptides of this invention are biologically active *mpl* ligand variant(s) that have an amino acid sequence having at least 70% amino acid sequence identity with the human *mpl* ligand (see Fig. 1 [SEQ ID NO: 1]), the murine *mpl* ligand (see Fig. 16 [SEQ ID NOS: 12 & 13]), the recombinant porcine *mpl* ligand (see Fig. 19 [SEQ ID NO: 18]) or the porcine *mpl* ligand isolated from aplastic porcine plasma, preferably at least 75%, more preferably at least 80%, still more preferably at least 85%, even more preferably at least 90%, and most preferably at least 95%.

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The mpl ligand isolated from aplastic porcine plasma has the following characteristics:

- (1) The partially purified ligand elutes from a gel filtration column run in either PBS, PBS containing 0.1% SDS or PBS containing 4M MgCl<sub>2</sub> with Mr of 60,000-70,000;
  - (2) The ligand's activity is destroyed by pronase;
  - (3) The ligand is stable to low pH (2.5), SDS to 0.1%, and 2M urea;
- (4) The ligand is a glycoprotein, based on its binding to a variety of lectin columns;
- 10 (5) The highly purified ligand elutes from non-reduced SDS-PAGE with a Mr of 25,000-35,000. Smaller amounts of activity also elute with Mr of ~18,000-22,000 and 60,000;
  - (6) The highly purified ligand resolves on reduced SDS-PAGE as a doublet with Mr of 28,000 and 31,000;
  - (7) The amino-terminal sequence of the 18,000-22,000, 28,000 and 31,000 bands is the same SPAPPACDPRLLNKLLRDDHVLHGR (SEQ ID NO: 29); and
    - (8) The ligand binds and elutes from the following affinity columns

Blue-Sepharose,

CM Blue-Sepharose,

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MONO-Q,

MONO-S.

Lentil lectin-Sepharose,

WGA-Sepharose,

Con A-Sepharose,

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Ether 650m Toyopearl,

Butyl 650 m Toyopearl,

Phenyl 650m Toyopearl, and

Phenyl-Sepharose.

More preferred *mpl* ligand polypeptides are those encoded by human genomic or cDNA having an amino acid sequence described in **Fig. 1** (SEQ ID NO: 1).

Other preferred naturally occurring biologically active *mpl* ligand polypeptides of this invention include prepro-*mpl* ligand, pro-*mpl* ligand, mature *mpl* ligand, *mpl* ligand fragments and glycosylation variants thereof.

Still other preferred polypeptides of this invention include *mpl* ligand sequence variants and chimeras. Ordinarily, preferred *mpl* ligand sequence variants and chimeras are biologically active *mpl* ligand variants that have an amino acid sequence having at least 70% amino acid sequence identity with the human *mpl* ligand or the *mpl* ligand isolated from aplastic porcine plasma, preferably at least 75%, more

preferably at least 80%, still more preferably at least 85%, even more preferably at least 90%, and most preferably at least 95%. An exemplary preferred *mpl* ligand variant is a N-terminal domain hML variant (refered to as the "EPO-domain" because of its sequence homology to erythropoietin). The prefered hML EPO-domain comprises about the first 153 amino acid residues of mature hML and is refered to as hML<sub>153</sub>. An optionally prefered hML sequence variant comprises one in which one or more of the basic or dibasic amino acid residue(s) in the C-terminal domain is substituted with a non-basic amino acid residue(s) (*e.g.*, hydrophobic, neutral, acidic, aromatic, Gly, Pro and the like). A prefered hML C-terminal domain sequence variant comprises one in which Arg residues 153 and 154 are replaced with Ala residues. This variant is refered to as hML<sub>332</sub>(R153A, R154A). An alternative prefered hML variant comprises either hML<sub>332</sub> or hML<sub>153</sub> in which amino residues 111-114 (QLPP or LPPQ) are deleted or replaced with a different tetrapeptide sequence(*e.g.* AGAG or the like). The foregoing deletion mutants are refered to as Δ4hML<sub>332</sub> or Δ4hML<sub>153</sub>.

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A preferred chimera is a fusion between *mpl* ligand or fragment (defined below) thereof with a heterologous polypeptide or fragment thereof. For example, hML<sub>153</sub> may be fused to an IgG fragment to improve serum half-life or to IL-3, G-CSF or EPO to produce a molecule with inhanced thrombopoietic or chimeric hematopoletic activity.

An alternative preferred human mpl ligand chimera is a "ML-EPO domain chimera" that consists of the N-terminus 153 to 157 hML residues substituted with one or more, but not all, of the human EPO residues approximately aligned as shown in Fig. 10 (SEQ ID NO: 7). In this embodiment, the hML chimera would be about 153-166 residues in length in which individual or blocks of residues from the human EPO sequence are added or substituted into the hML sequence at positions corresponding to the alignment shown in Fig. 10 (SEQ ID NO: 6). Exemplary block sequence inserts into the N-terminus portion of hML would include one or more of the N-glycosylation sites at positions (EPO) 24-27, 38-40, and 83-85; one or more of the four predicted amphipathic α-helical bundles at positions (EPO) 9-22, 59-76, 90-107, and 132-152; and other highly conserved regions including the N-terminus and Cterminus regions and residue positions (epo) 44-52 (see e.g., Wen et al., Blood, 82:1507-1516 [1993] and Boissel et al., J. Biol. Chem., 268(21):15983-15993 It is contemplated this "ML-EPO domain chimera" will have mixed (1993]). thrombopoietic-erythropoletic (TEPO) biological activity.

Other preferred polypeptides of this invention include *mpl* ligand fragments having a consecutive sequence of at least 10, 15, 20, 25, 30, or 40 amino acid residues that are identical to the sequences of the *mpl* ligand isolated from aplastic

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porcine plasma or the human *mpl* ligand described herein (see *e.g.* **Table 14**, **Example 24**). A preferred *mpl* ligand fragment is human ML[1-X] where X is 153, 164, 191, 205, 207, 217, 229, or 245 (see Fig. 1 [SEQ ID NO: 1] for the sequence of residues 1-X). Other preferred *mpl* ligand fragments include those produced as a result of chemical or enzymatic hydrolysis or digestion of the purified ligand.

Another preferred aspect of the invention is a method for purifying *mpl* ligand molecules comprises contacting a *mpl* ligand source containing the *mpl* ligand molecules with an immobilized receptor polypeptide, specifically *mpl* or a *mpl* fusion polypeptide, under conditions whereby the *mpl* ligand molecules to be purified are selectively adsorbed onto the immobilized receptor polypeptide, washing the immobilized support to remove non-adsorbed material, and eluting the molecules to be purified from the immobilized receptor polypeptide with an elution buffer. The source containing the *mpl* ligand may be plasma where the immobilized receptor is preferably a *mpl*-lgG fusion.

Alternatively, the source containing the mpl ligand is recombinant cell culture where the concentration of mpl ligand in either the culture medium or in cell lysates is generally higher than in plasma or other natural sources. In this case the above described mpl-lgG immunoaffinity method, while still useful, is usually not necessary and more traditional protein purification methods known in the art may be applied. Briefly, the preferred purification method to provide substantially homogeneous mpl ligand comprises: removing particulate debris, either host cells or lysed fragments by, for example, centrifugation or ultrafiltration; optionally, protien may be concentrated with a commercially available protein concentration filter; followed by separating the ligand from other impurities by one or more steps selected from; immunoaffinity, ion-exchange (e.g., DEAE or matricles containing carboxymethyl or sulfopropyl groups), Blue-Sepharose, CM Blue-Sepharose, MONO-Q, MONO-S, lentil lectin-Sepharose, WGA-Sepharose, Con A-Sepharose, Ether Toypearl, Butyl Toypearl, Phenyl Toypearl, protein A Sepharose, SDS-PAGE, reverse phase HPLC (e.g., silica gel with appended aliphatic groups) or Sephadex molecular seive or size exclusion chromatography, and ethanol or ammonium sulfate precipitation. A protease inhibitor such as methylsulfonylfluoride (PMSF) may be included in any of the foregoing steps to inhibit proteolysis.

In another preferred embodiment, this invention provides an isolated antibody capable of binding to the *mpl* ligand. A preferred *mpl* ligand isolated antibody is monoclonal (Kohler and Milstein, *Nature*, 256:495-497 [1975]; Campbell, *Laboratory Techniques in Biochemistry and Molecular Biology*, Burdon *et al.*, Eds, Volume 13, Elsevier Science Publisrers, Amsterdam [1985]; and Huse *et al.*, *Science*, 246:1275-1281 [1989]). Preferred *mpl* ligand isolated antibody is one that blnds

to mpl ligand with an affinity of at least about  $10^6$  l/mole. More preferably the antibody blnds with an affinity of at least about  $10^7$  l/mole. Most preferably, the antibody is raised against the mpl ligand having one of the above described effector functions. The isolated antibody capable of binding to the mpl ligand may optionally be fused to a second polypeptide and the antibody or fusion thereof may be used to isolate and purify mpl ligand from a source as described above for immobilized mpl polypeptide. In a further preferred aspect of this embodiment, the invention provides a method for detecting the mpl ligand in vitro or in vivo comprising contacting the antibody with a sample, especially a serum sample, suspected of containing the ligand and detecting if binding has occurred.

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In still further preferred embodiments, the invention provides an isolated nucleic acid molecule encoding the mpl ligand or fragments thereof, which nucleic acid molecule may be labeled or unlabeled with a detectable moiety, and a nucleic acid molecule having a sequence that is complementary to, or hybridizes under stringent or moderately stringent conditions with, a nucleic acid molecule having a sequence encoding a mpl ligand. A preferred mpl ligand nucleic acid is RNA or DNA that encodes a biologically active mpl ligand sharing at least 75% sequence identity, more preferably at least 80%, still more preferably at least 85%, even more preferably 90%, and most preferably 95% sequence identity with the human mpl ligand. More preferred isolated nucleic acid molecules are DNA sequences encoding biologically active mpl ligand, selected from: (a) DNA based on the coding region of a mammalian mpl ligand gene (e.g., DNA comprising the nucleotide sequence provided in Fig. 1 (SEQ ID NO: 2), or fragments thereof); (b) DNA capable of hybridizing to a DNA of (a) under at least moderately stringent conditions; and (c) DNA that is degenerate to a DNA defined in (a) or (b) which results from degeneracy of the genetic code. It is contemplated that the novel mpl ligands described herein may be members of a family of ligands or cytokines having suitable sequence identity that their DNA may hybridize with the DNA of Fig. 1 (SEQ ID NO: 2) (or the complement or fragments thereof) under low to moderate stringency conditions. Thus a further aspect of this invention includes DNA that hybridizes under low to moderate stringency conditions with DNA encoding the mpl ligand polypeptides.

In a further preferred embodiment of this invention, the nucleic acid molecule is cDNA encoding the *mpl* ligand and further comprises a replicable vector in which the cDNA is operably linked to control sequences recognized by a host transformed with the vector. This aspect further includes host cells transformed with the vector and a method of using the cDNA to effect production of *mpl* ligand, comprising expressing the cDNA encoding the *mpl* ligand in a culture of the transformed host cells and recovering the *mpl* ligand from the host cell culture. The *mpl* ligand prepared in this manner is

preferably substantially homogeneous human mpl ligand. A preferred host cell for producing mpl ligand is Chinese hamster ovary (CHO) cells.

The invention further includes a preferred method for treating a mammal having an immunological or hematopoietic disorder, especially thrombocytopenia comprising administering a therapeutically effective amount of a *mpl* ligand to the mammal. Optionally, the *mpl* ligand is administered in combination with a cytokine, especially a colony stimulating factor or interleukin. Preferred colony stimulating factors or interleukins include; kit-ligand, LIF, G-CSF, GM-CSF, M-CSF, EPO, IL-1, IL-2, IL-3, IL-5, IL-6, IL-7, IL-8, IL-9 or IL-11.

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### III. Methods of Making

Platelet production has long been thought by some authors to be controlled by multiple lineage specific humoral factors. It has been postulated that two distinct cytokine activities, referred to as megakaryocyte colony-stimulating factor (meg-CSF) and thrombopoietin, regulate megakaryocytopoiesis and thrombopoiesis (Williams et al., J. Cell Physiol., 110:101-104 [1982]; Williams et al., Blood Cells, 15:123-133 [1989]; and Gordon et al., Blood, 80:302-307 [1992]). According to this hypothesis, meg-CSF stimulates the proliferation of progenitor megakaryocytes while thrombopoletin primarily affects maturation of more differentiated cells and ultimately platelet release. Since the 1960's the induction and appearance of both meg-CSF and thrombopoietin activities in the plasma, serum and urine of animals and humans following thrombocytopenic episodes has been well documented (Odell et al., Proc. Soc. Exp. Biol. Med., 108:428-431 [1961]; Nakeff et al., Acta Haematol., 54:340-344 [1975]; Specter, Proc. Soc. Exp. Biol., 108:146-149 [1961]; Schreiner et al., J. Clin. Invest., 49:1709-1713 [1970]; Ebbe, Blood, 44:605-608 [1974]; Hoffman et al., N. Engl. J. Med., 305:533 [1981]; Straneva et al., Exp. Hematol., 17:1122-1127 [1988]; Mazur et al., Exp. Hematol., 13:1164 [1985]; Mazur et al., J.Clin. Invest., 68:733-741 [1981]; Sheiner et al., Blood, 56:183-188 [1980]; Hill et al., Exp. Hematol., 20:354-360 [1992]; and Hegyi et al., Int. J. Cell Cloning, 8:236-244 [1990]). These activities were reported to be lineage specific and distinct from known cytokines (Hill R.J. et al., Blood 80:346 (1992); Erickson-Miller C.L. et al., Brit. J. Haematol., 84:197-203 (1993); Straneva J.E. et al., Exp. Hematol. 20:4750(1992); and Tsukada J. et al., Blood 81:866-867 [1993]). Heretofore, attempts to purify meg-CSF or thrombopoietin from thrombocytopenic plasma or urine have been unsuccessful.

Consistent with the above observations describing thrombocytopenic plasma, we have found that aplastic porcine plasma (APP) obtained from irradiated pigs stimulates human megakaryocytopolesis in vitro. We have found that this stimulatory

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activity is abrogated by the soluble extracellular domain of c-mpl, confirming APP as a potential source of the putative mpl ligand (ML). We have now successfully purified the mpl ligand from APP and amino acid sequence information was used to isolate murine, porcine and human ML cDNA. These ML's have sequence homology to erythropoietin and have both meg-CSF and thrombopoietin-like activities.

### 1. Purification and Identification of mpl Ligand from Plasma

As set forth above, aplastic plasma from a variety of species has been reported to contain activities that stimulate hematopolesis *in vitro*, however no hematopoletic stimulatory factor has previously been reported isolated from plasma. One source of aplastic plasma is that obtained from irradiated pigs. This aplastic porcine plasma (APP) stimulates human hematopolesis *in vitro*. To determine if APP contained the *mpl* ligand, its effect was assayed by measuring <sup>3</sup>H-thymidine incorporation into Ba/F3 cells transfected with human *mpl* P (Ba/F3-*mpl*) by the procedure shown in Fig. 2. APP stimulated <sup>3</sup>H-thymidine incorporation into Ba/F3-*mpl* cells but not Ba/F3 control cells (*i.e.*, not transfected with human *mpl* P). Additionally, no such activity was observed in normal porcine plasma. These results indicated that APP contained a factor or factors that transduced a proliferative signal through the *mpl* receptor and therefore might be the natural ligand for this receptor. This was futher supported by the finding that treatment of APP with soluble *mpl*-IgG blocked the stimulatory effects of APP on Ba/F3-*mpl* cells.

The activity in APP appeared to be a protein since pronase, DTT, or heat destroy the activity in APP (Fig. 3). The activity was also non-dialyzable. The activity was, however, stable to low pH (pH 2.5 for 2 hrs.) and was shown to bind and elute from several lectin-affinity columns, indicating that it was a glycoprotein. To further elucidate the structure and identity of this activity it was affinity purified from APP using a *mpl*-IgG chimera.

APP was treated according to the protocol set forth in **Examples 1** and **2**. Briefly, the *mpl* ligand was purified using hydrophobic interaction chromatography (HIC), immobilized dye chromatography, and *mpl*-affinity chromatography. The recovery of activity from each step is shown in Fig. 4 and the fold purification is provided in **Table 1**. The overall recovery of activity through the *mpl*-affinity column was approximately 10%. The peak activity fraction (F6) from the *mpl*-affinity column has an estimated specific activity of 9.8 x10<sup>6</sup> units/mg. The overall purification from 5 liters of APP was approximately 4 x10<sup>6</sup> fold (0.8 units/mg to 3.3 x 10<sup>6</sup> units/mg) with a 83 x 10<sup>6</sup>-fold reduction in protein (250 gms to 3  $\mu$ g). We estimated the specific activity of the ligand eluted from the *mpl*-affinity column to be ~3x10<sup>6</sup> units/mg.

TABLE 1

Purification of mpl Ligand

		Full	ilcation (	<u>,, ,,, ,, , , , , , , , , , , , , , , </u>			
Sample	Volume	Protein	Units/ml	Units	Specific Acitivity	Yield	Fold
Campio	mls	mg/ml			Units/mg	%	Purification
APP	5000	50	40	200,000	0.8	_	1
Phenyl	4700	0.8	40	200,000	50	94	62
Blue-Sep.	640	0.93	400	256,000	430	128	538
mpl (μ1)							
(Fxns 5-7)	12_	5x10-4	1666	20,000	3,300,000	10	4,100,000

Protein was determined by the Bradford assay. Protein concentration of *mpl*-eluted fractions 5-7 are estimates based on staining intensity of a silver stained SDS-gel. One unit is defined as that causing 50% maximal stimulation of Ba/F3-*mpl* cell proliferation.

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Analysis of eluted fractions from the *mpl* affinity column by SDS-PAGE (4-20%, Novex gel) run under reducing conditions, revealed the presence of several proteins (Fig. 5). Proteins that silver stained with the strongest intensity resolved with apparent Mr of 66,000, 55,000, 30,000, 28,000 and 18,000-22,000. To determine which of these proteins stimulated proliferation of Ba/F3-*mpl* cell cultures, the proteins were eluted from the gel as described in Example 2.

The results of this experiment showed that most of the activity eluted from a get slice that included proteins with Mr 28,000-32,000, with lesser activity eluting in the 18,000-22,000 region of the get (Fig. 6). The only proteins visible in these regions had Mr of 30,000, 28,000 and 18,000-22,000. To Identify and obtain protein sequence for the proteins resolving in this region of the get (i.e. bands at 30, 28 and 18-22 kDa), these three proteins were electroblotted to PVDF and sequenced as described in Example 3. Amino-terminus sequences obtained are provided in Table 2.

TABLE 2

Mpi Ligand Amino-Terminus Sequences

30 kDa						
1	5	10	15	20	25	
(S) P A	PPA(C	<u>)DPRLI</u>	NKLLR	D D (H/S) V L	H (G) R L	(SEQ ID NO: 30)
28 kDa						
1	5	10	15	20	25	
(S) P A	PPAX	DPRLL	NKLLR	D D (H) V L (H)	GR	(SEQ ID NO: 31)
18-22	kDa					
1	5	10				
XPA	PPAX	DPRLX	(N) (K)			(SEQ ID NO: 32)

Computer-assisted analysis revealed these amino acid sequences to be novel. Because all three sequences were the same, it was believed the 30 kDa, 28 kDa and 18-22 kDa proteins were related and might be different forms of the same novel protein. Futhermore, this protein(s) was a likely candidate as the natural *mpl* ligand because the activity resolved on SDS-PAGE in the same region (28,000-32,000) of a 4-20% gel. In addition, the partially purified ligand migrated with a Mr of 17,000-30,000 when subjected to gel filtration chromatography using a Superose 12 (Pharmacia) column. It is believed the different Mr forms of the ligand are a result of proteolysis or glycosylation differences or other post or pre-translational modifications.

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As described earlier, antisense human mpl RNA abrogated megakaryocytopolesis in human bone marrow cultures enriched with CD 34+ progenitor cells without affecting the differentiation of other hematopoietic cell lineages (Methia et al., supra). This result suggested that the mpl receptor might play a role in the differentiation and proliferation of megakaryocytes in vitro. To further elucidate the role of the mpl ligand in megakaryocytopolesis, the effects of APP and mpl ligand depleted APP on in vitro human megakaryocytopolesis was compared. The effect of APP on human megakaryocytopoiesis was determined using a modification of the liquid suspension megakaryocytopoiesis assay described in Example 4. In this assay, human peripheral stem cells (PSC) were treated with APP before and after mpl-lgG affinity chromatography. GPIIbIIIa stimulation of megakaryocytopolesis was quantitated with an  $^{125}$ I-anti-II<sub>b</sub>III<sub>a</sub> antibody (Fig. 7). Shown in Fig. 7, 10% APP caused approximately a 3-fold stimulation while APP depleted of mpl ligand had no effect. Significantly, the mpl ligand depleted APP did not induce proliferation of the Ba/F3-mp/ cells.

In another experiment, soluble human *mpl*-IgG added at days 0, 2 and 4 to cultures containing 10% APP neutralized the stimulatory effects of APP on human megakaryocytopoiesis (**Fig. 8**). These results indicate that the *mpl* ligand plays a role in regulating human megakaryocytopoiesis and therefore may be useful for the treatment of thrombocytopenia.

# 2. Molecular Cloning of the mpl Ligand

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Based on the amino-terminal amino acid sequence obtained from the 30 kDa, 28 kDa and 18-22 kDa proteins (see **Table 2** above), two degenerate oligonucleotide primer pools were designed and used to amplify porcine genomic DNA by PCR. It was reasoned that If the amino-terminal amino acid sequence was encoded by a single exon then the correct PCR product was expected to be 69 bp long. A DNA fragment of this size was found and subcloned into pGEMT. The sequences of the oligonucleotide PCR primers and the three clones obtained are shown in **Example 5**. The amino acid sequence (PRLLNKLLR [SEQ ID NO: 33]) of the peptide encoded between the PCR primers was identical to that obtained by amino-terminal protein sequencing of the porcine ligand (see residues 9-17 for the 28 and 30 kDa porcine protein sequences above).

A synthetic oligonucleotide based on the sequence of the PCR fragment was used to screen a human genomic DNA library. A 45-mer oligonucleotide, designated pR45, was designed and synthesized based on the sequence of the PCR fragment. This oligonucleotide had the following sequence:

# 5' GCC-GTG-AAG-GAC-GTG-GTC-GTC-ACG-AAG-CAG-TTT-ATT-TAG-GAG-TCG 3' (SEQ ID NO: 34)

This deoxyoligonucleotide was used to screen a human genomic DNA library in \$\lambda gem12\$ under low stringency hybridization and wash conditions according to **Example 6.** Positive clones were picked, plaque purified and analyzed by restriction mapping and southern blotting. A 390 bp EcoRI-Xbal fragment that hybridized to the 45-mer was subcloned into pBluescript SK-. DNA sequencing of this clone confirmed that DNA encoding the human homolog of the porcine *mpl* ligand had been isolated. The human DNA sequence and deduced amino acid sequence are shown in **Fig. 9** (SEQ ID NOS: 3 & 4). The predicted positions of introns in the genomic sequence are also indicated by arrows, and define a putative exon ("exon 3").

Based on the human "exon 3" sequence (Example 6) oligonucleotides corresponding to the 3' and 5' ends of the exon sequence were synthesized. These 2 primers were used in PCR reactions employing as a template cDNA prepared from various human tissues. The expected size of the correct PCR product was 140 bp. After analysis of the PCR products on a 12% polyacrylamide gel, a DNA fragment of the

expected size was detected in cDNA libraries prepared from human adult kidney, 293 fetal kidney cells and cDNA prepared from human fetal liver.

A fetal liver cDNA library (7x10<sup>6</sup> clones) in lambda DR2 was next screened with the same 45-mer oligonucleotide used to screen the human genomic library and the fetal liver cDNA library under low stringency hybridization conditions. Positive clones were picked, plaque purified and the insert size was determined by PCR. One clone with a 1.8 kb insert was selected for further analysis. Using the procedures described in **Example 7** the nucleotide and deduced amino acid sequence of the human *mpl* ligand (hML) were obtained. These sequences are presented in **Fig. 1** (SEQ ID NOS: 1 & 2).

### 3. Structure of the Human mpl Ligand (hML)

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The human *mpl* ligand (hML) cDNA sequence (Fig. 1 [SEQ ID NO: 2]) comprises 1774 nucleotides followed by a poly(A) tail. It contains 215 nucleotides of 5' untranslated sequence and a 3' untranslated region of 498 nucleotides. The presumed initiation codon at nucleotide position (216-218) is within a consensus sequence favorable for eukaryotic translation initiation. The open reading frame is 1059 nucleotides long and encodes a 353 amino acid residue polypeptide, beginning at nucleotide position 220. The N-terminus of the predicted amino acid sequence is highly hydrophobic and probably corresponds to a signal peptide. Computer analysis of the predicted amino acid sequence (von Heijne *et al.*, *Eur. J. Biochem.*, 133:17-21 [1983]) indicates a potential cleavage site for signal peptidase between residues 21 and 22. Cleavage at that position would generate a mature polypeptide of 332 amino acid residues beginning with the amino-terminal sequence obtained from *mpl* ligand purified from porcine plasma. The predicted non-glycosylated molecular weight of the 332 amino acid residue ligand is about 38 kDa. There are 6 potential N-glycosylation sites and 4 cysteine residues.

Comparison of the *mpl* ligand sequence with the Genbank sequence database revealed 23% identity between the amino terminal 153 residues of mature human *mpl* ligand and human erythropoietin (Fig. 10 [SEQ ID NOS: 6 & 7]). When conservative substitutions are taken into account, this region of hML shows 50% similarity to human erythropoietin (hEPO). Both hEPO and the hML contain four cysteines. Three of the 4 cysteines are conserved in hML, including the first and last cysteines. Site-directed mutagenesis experiments have shown that the first and last cysteines of erythropoietin form a disulfide bond that is required for function (Wang, F.F.et al., Endocrinology 116:2286-2292 [1983]). By analogy, the first and last cysteines of hML may also form a critical disulfide bond. None of the glycosylation sites are

conserved in hML. All potential hML N-linked glycosylation sites are located in the carboxy-terminal half of the hML polypeptide.

Similar to hEPO, the hML mRNA does not contain the consensus polyadenylation sequence AAUAAA, nor the regulatory element AUUUA that is present in 3' untranslated regions of many cytokines and is thought to influence mRNA stability (Shawet al., Cell, 46:659-667 [1986]). Northern blot analysis reveals low levels of a single 1.8 kb hML RNA transcript in both fetal and adult liver. After longer exposure, a weaker band of the same size could be detected in adult kidney. By comparison, human erythropoietin is expressed in fetal liver and, in response to hypoxia, the adult kidney and liver (Jacobs et al., Nature, 313:804-809 [1985] and Bondurant et al., Molec. Cell. Biol., 6:2731-2733 [1986]).

The importance of the C-terminal region of the hML remains to be elucidated. Based on the presence of the six potential sites for N-linked glycosylation and the ability of the ligand to bind lectin-affinity columns, this region of the hML is likely glycosylated. In some gel elution experiments, we observed activity resolving with a M<sub>r</sub> around 60,000 which may represent the full length, glycosylated molecule. The C-terminal region may therefore act to stabilize and increase the half-life of circulating hML. In the case of erythropoletin, the non-glycosylated form has full in vitro biological activity, but has a significantly reduced plasma half-life relative to glycosylated erythropoietin (Takeuchi et al., J. Biol. Chem., 265:12127-12130 [1990]; Narhi et al., J. Biol. Chem., 266:23022-23026 [1991] and Spivack et al., Blood, 7:90-99 [1989]). The C-terminal domain of hML contains two di-basic amino acid sequences [Arg-Arg motifs at positions 153-154 and 245-246] that could serve as potential processing sites. Cleavage at these sites may be responsible for generating the 30, 28 and 18-22 kDa forms of the ML isolated from APP. Significantly, the Arg<sub>153</sub>-Arg<sub>154</sub> sequence occurs immediately following the erythropoietin-like domain of the ML. These observations indicate that full length ML may represent a precursor protein that undergoes limited proteolysis to generate the mature ligand.

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# 4. Isoforms and Variants of the Human mpl Ligand

Isoforms or alternatively spliced forms of human *mpl* ligand were detected by PCR in human adult liver. Briefly, primers were synthesized corresponding to each end as well as selected internal regions of the coding sequence of hML. These primers were used in RT-PCR to amplify human adult liver RNA as described in **Example 10**. In addition to the full length form, designated hML, three other forms, designated hML2, hML3 and hML4, were observed or deduced. The mature deduced amino acid sequences of all four isoforms is presented in **Fig. 11** (SEQ ID NOS: 6, 8, 9 & 10).

hML3 has a 116 nucleotide deletion a position 700 which results in both an amino acid deletion and a frameshift. The cDNA now encodes a mature polypeptide that is 265 amino acid long and diverges from the hML sequence at amino acid residue 139. Finally, hML4 has both a 12 nucleotide deletion following nucleotide position 618 (also found in the mouse and the pig sequences [see below]) and the 116 bp deletion found in hML3. Altough no clones with only the 12 bp deletion (following nucleotide 619) have been isolated in the human (designated hML2), this form is likely to exist because such a isoform has been identified in both the mouse and pig (see below), and because it has been identified in conjunction with the 116 nucleotide deletion in hML4.

Both a substitutional variant of hML in which the dibasic Arg<sub>153</sub>-Arg<sub>154</sub> sequence was replaced with two alanine residues and a "EPO-domain" truncated form of hML were constructed to determine whether the full length ML was necessary for biological activity. The Arg<sub>153</sub>-Arg<sub>154</sub> dibasic sequence substitutional variant, referred to as hML(R153A, R154A), was constructed using PCR as described in **Example 10**. The "EPO-domain" truncated form, hML<sub>153</sub>, was also made using PCR by introducing a stop codon following Arg153.

# Expression of Recombinant Human mpl Ligand (rhML) in Transiently Transfected Human Embryonic Kidney (293) Cells

To confirm that the cloned human cDNA encoded a ligand for *mpl*, the ligand was expressed in mammalian 293 cells under the control of the cytomegalovirus immediate early promoter using the expression vectors pRK5-hML or pRK5-hML153. Supernatants from transiently transfected human embryonic kidney 293 cells were found to stimulate <sup>3</sup>H-thymidine incorporation in Ba/F3-*mpl* cells, but not in parental Ba/F3 cells (Fig. 12A). Media from the 293 cells transfected with the pRK vector alone did not contain this activity. Addition of *mpl*-lgG to the media abolished the stimulation (data not shown). These results show that the cloned cDNA encodes a functional human ML (hML).

To determine if the "EPO-domain" alone could bind and activate mpl, the truncated form of hML, rhML<sub>153</sub>, was expressed in 293 cells. Supernatants from transfected cells were found to have activity similar to that present in supernatants from cells expressing the full length hML (Fig. 12A), indicating that the C-terminal domain of ML is not required for binding and activation of c-mpl.

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# 6. mpl Ligand Stimulates Megakaryocytopolesis and Thrombopolesis

Both the full length rhML and the truncated rhML<sub>153</sub> forms of recombinant hML stimulated human megakaryocytopoiesis *in vitro* (Fig. 12B). This effect was observed in the absence of other exogenously added hematopoietic growth factors. With the exception of IL-3, the ML was the only hematopoietic growth factor tested that exhibited this activity. IL-11, IL-6, IL-1, erythropoietin, G-CSF, IL-9, LIF, kit ligand (KL), M-CSF, OSM and GM-CSF had no effect on megakaryocytopoiesis when tested separately in our assay (data not shown). This result demonstrates that the ML has megakaryocyte-stimulating activity, and indicates a role for ML in regulating megakaryocytopoiesis.

Thrombopoietic activities present in plasma of thrombocytopenic animals have been shown to stimulate platelet production in a mouse rebound thrombocytosis assay (McDonald, Proc. Soc. Exp. Biol. Med., 14:1006-1001 [1973] and McDonald et al., Scand. J. Haematol., 16:326-334 [1976]). In this model mice are made acutely thrombocytopenic using specific antiplatelet serum, resulting in a predictable rebound Such immuno-thrombocythemic mice are more responsive to thrombocytosis. exogenous thrombopoietin-like activities than are normal mice (McDonald, Proc. Soc. Exp. Biol. Med., 14:1006-1001 [1973]), just as exhypoxic mice are more sensitive to erythropoietin than normal are mice (McDonald, et al., J. Lab. Clin. Med., 77:134-143 [1971]). To determine whether the rML stimulates platelet production in vivo, mice in rebound thrombocytosis were injected with partially purified rhML. Platelet counts and incorporation of 35S into platelets were then quantitated. Injection of mice with 64,000 or 32,000 units of rML significantly increased platelet production, as evidenced by a ~20% increase in platelet counts (p=0.0005 and 0.0001, respectively) and a ~40% increase in 35S incorporation into platelets (p=0.003) in the treated mice versus control mice injected with excipient alone (Fig. 12C). This level of stimulation is comparable to that which we have observed with IL-6 in this model (data not shown). Treatment with 16,000 units of rML did not significantly stimulate platelet production. These results indicate that ML stimulates platelet production in a dose-dependent manner and therefore possesses thrombopoietin-like activity.

293 cells were also transfected with the other hML isoform constructs described above and the supernatants were assayed using the Ba/F3-mpl proliferation assay (see Fig. 13). hML2 and hML3 showed no detectable activity in this assay, however the activity of hML(R153A, R154A) was similar to hML and hML153 indicating that processing at the Arg153-Arg154 di-basic site is neither required for nor detrimental to activity.

### 7. Megakaryocytopoiesis and the mpl Ligand

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It has been proposed that megakaryocytopolesis is regulated at multiple cellular levels (Williams et al., J.Cell Physiol., 110:101-104 [1982] and Williams et al., Blood Cells, 15:123-133 [1989]). This is based largely on the observation that certain hematopoietic growth factors stimulate proliferation of megakaryocyte progenitors while others appear to primarily affect maturation. The results presented here suggest that the ML acts both as a proliferative and maturation factor. That ML stimulates proliferation of megakaryocyte progenitors is supported by several lines of First, APP stimulates both proliferation and maturation of human evidence. megakaryocytes in vitro, and this stimulation is completely inhibited by mpl-IgG (Figs. 7 and 8). Furthermore, the inhibition of megakaryocyte colony formation by c-mpl antisense oligonucleotides (Methia et al., Blood, 82:1395-1401 [1993]) and the finding that c-mpl can transduce a proliferative signal in cells into which it is transfected (Skoda et al., EMBO, 12:2645-2653 [1993] and Vigon et al., Oncogene, 8:2607-2615 [1993]) also indicate that ML stimulates proliferation. The apparent expression of c-mpl during all stages of megakaryocyte differentiation (Methia et al., Blood, 82:1395-1401 [1993]) and the ability of recombinant ML to rapidly stimulate platelet production in vivo indicate that ML also affects maturation. The availability of recombinant ML makes possible a careful evaluation of its role in regulating megakaryocytopoiesis and thrombopoiesis as well as its potential to influence other hematopoietic lineages.

### 8. Isolation of the Human mpi Ligand (TPO) Gene

Human genomic DNA clones of the TPO gene were isolated by screening a human genomic library in  $\lambda$ -Gem12 with pR45, under low stringency conditions or under high stringency conditions with a fragment corresponding to the 3' half of human cDNA coding for the mpl ligand. Two overlapping lambda clones spanning 35 kb were isolated. Two overlapping fragments (BamH1 and EcoRl) containing the entire TPO gene were subcloned and sequenced (see **Figs. 14A, 14B** and **14C**).

The structure of the human gene is composed of 6 exons within 7 kb of genomic DNA. The boundaries of all exon/intron junctions are consistent with the consensus motif established for mammalian genes (Shapiro, M. B., et al., Nucl. Acids Res. 15:7155 [1987]). Exon 1 and exon 2 contain 5' untranslated sequence and the initial four amino acids of the signal peptide. The remainder of the secretory signal and the first 26 amino acids of the mature protein are encoded within exon 3. The entire carboxyl domain and 3' untranslated as well as ~50 amino acids of the erythropoletin-

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like domain are encoded within exon 6. The four amino acids involved in the deletion observed within hML-2 (hTPO-2) are encoded at the 5' end of exon 6.

Analysis of human genomic DNA by Southern blot indicated the gene for TPO is present in a single copy. The chromosomal location of the gene was determined by fluorescent in situ hybridization (FISH) which mapped to chromosome 3q27-28.

# 9. Expression and Purification of TPO from 293 Cells

Preperation and purification of ML or TPO from 293 cells is described in detail in **Example 19**. Briefly, cDNA corresponding to the TPO entire open reading frame was obtained by PCR using pRK5-hmpl I. The PCR product was purified and cloned between the restriction sites Clal and Xbal of the plasmid pRK5tkneo (a pRK5 derived vector modified to express a neomycin resistance gene under the control of the thymidine kinase promote) to obtain the vector pRK5tkneo.ORF(a vector coding for the entire open reading frame).

A second vector coding for the EPO homologous domain was generated the same but using different PCR primers to obtain the final construct called pRK5-tkneoEPO-D.

These two constructs were transfected into Human Embryonic Kidney cells by the CaPO<sub>4</sub> method and neomycin resistant clones were selected and allowed to grow to confluency. Expression of ML<sub>153</sub> or ML<sub>332</sub> in the conditioned media from these clones was assessed using the Ba/F3-mpl proliferation assay.

Purification of rhML332 was conducted as described in Example 19. Briefly, 293-rhML332 conditioned media was applied to a Blue-Sepharose (Pharmacia) column that was subsequently washed with a buffer containing 2M urea. The column was eluted with a buffer containing 2M urea and 1M NaCl. The Blue-Sepharose elution pool was then directly applied to a WGA-Sepharose column, washed with 10 column volumes of buffer containing 2M urea and 1 M NaCl and eluted with the same buffer containing 0.5M N-acetyl-D-glucosamine. The WGA-Sepharose eluate was applied to a C4-HPLC column (Synchrom, Inc.) and eluted with a discontinuous propanol gradient. By SDS-PAGE the purified 293-rhML332 migrates as a broad band in the 68-80 kDa region of the gel (see Fig. 15).

Purification of rhML<sub>153</sub> was also conducted as described in **Example 19**. Briefly, 293-rhML<sub>153</sub> conditioned media was resolved on Blue-Sepharose as described for rhML<sub>332</sub>. The Blue Sepharose eluate was applied directly to a *mpl*-affinity column as described above. RhML<sub>153</sub> eluted from the *mpl*-affinity column was purified to homogeneity using a C4-HPLC column run under the same conditions used for rhML<sub>332</sub>. By SDS-PAGE the purified rhML<sub>153</sub> resolves into 2 major and 2 minor bands with Mr of ~18,000-22,000 (see **Fig. 15**).

### 10. The Murine mpl Ligand

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A DNA fragment corresponding to the coding region of the human mpl ligand was obtained by PCR, gel purified and labeled in the presence of 32P-dATP and 32P-dCTP. This probe was used to screen 106 clones of a mouse liver cDNA library in λGT10. A murine clone (Fig. 16 [SEQ ID NOS: 12 & 13]) containing a 1443 base pair insert was isolated and sequenced. The presumed initiation codon at nucleotide position 138-141 was within a consensus sequence favorable for eukaryotic translation initiation (Kozak, M. J.Cell Biol., 108:229-241 [1989]). This sequence defines an open reading frame of 1056 nucleotides, which predicts a primary translation product of 352 amino acids. Flanking this open reading frame are 137 nucleotides of 5' and 247 nucleotides of 3' untranslated sequence. There is no poly(A) tail following the 3' untranslated region indicating that the clone is probably not complete. The N-terminus of the predicted amino acid sequence is highly hydrophobic and probably represents a signal peptide. Computer analysis (von Heijne, G. Eur. J. Biochem. 133:17-21 [1983]) indicated a potential cleavage site for signal peptidase between residues 21 and 22. Cleavage at that position would generate a mature polypeptide of 331 amino acids (35 kDa) identified as mML331 (or mML2 for reasons described below). The sequence contains 4 cysteines, all conserved in the human sequence, and seven potential N-glycosylation sites, 5 of which are conserved in the human sequence. Again, as with hML, all seven potential N-glycosylation sites are located in the Cterminal half of the protein.

When compared with the human ML, considerable identity for both nucleotide and deduced amino acid sequences were observed in the "EPO-domains" of these ML's. However, when deduced amino acid sequences of human and mouse ML's were aligned, the mouse sequence appeared to have a tetrapeptide deletion between residues 111-114 corresponding to the 12 nucleotide deletion following nucleotide position 618 seen in both the human (see above) and pig (see below) cDNA's. Accordingly, additional clones were examined to detect possible murine ML isoforms. One clone encoded a 335 amino acid deduced sequence polypeptide containing the "missing" tetrapeptide LPLQ. This form is believed to be the full length murine ML and is refered to as mML or mML335. The nucleotide and deduced amino acid sequence for mML are provided in Fig. 17 (SEQ ID NOS: 14 & 15). This cDNA clone consists of 1443 base pairs followed by a poly(A) tail. It possesses an open reading frame of 1068 bp flanked by 134 bases of 5' and 241 bases of 3' untranslated sequence. The presumed initiation codon lies at nucleotide position 138-140. The open reading frame encodes a predicted protein of 356 amino acids, the first 21 of which are highly hydrophobic and likely function as a secretion signal.

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Finally, a third murine clone was isolated, sequenced and was found to contained the 116 nucleotide deletion corresponding to hML3. This murine isoform is therefore denominated mML3. Comparison of the deduced amino acid sequences of these two isoforms is shown in Fig. 18 (SEQ ID NOS: 9 & 16).

The overall amino acid sequence identity between human and mouse ML (Fig. 19 [SEQ ID NOS: 6 & 17]) is 72% but this homology is not evenly distributed. The region defined as the "EPO-domain" (amino acids 1-153 for the human sequence and 1-149 for the mouse) is better conserved (86% homology) than the carboxy-terminal region of the protein (62% homology). This may further indicate that only the "EPO-domain" is important for the biological activity of the protein. Interestingly, of the two di-basic amino acid motifs found in hML, only the di-basic motif immediately following the "EPO-domain" (residue position 153-154) in the human sequence is present in the murine sequence. This is consistent with the possibility that the full length ML may represent a precursor protein that undergoes limited proteolysis to generate the mature ligand. Alternatively, proteolysis between Arg153-Arg154 may facilitate hML clearance

An expression vector containing the entire coding sequence of mML was transiently transfected into 293 cells as described in **Example 1**. Conditioned media from these cells stimulated <sup>3</sup>H-thymidine incorporation into Ba/F3 cells expressing either murine or human *mpl* but had no effect on the parental (*mpl*-less) cell line. This indicates that the cloned murine ML cDNA encodes a functional ligand that is able to activate both the murine and human ML receptor (*mpl*).

## 11. The Porcine mpl Ligand

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Porcine ML (pML) cDNA was isolated by RACE PCR as described in **Example** 13. A PCR cDNA product of 1342 bp was found in kidney and subcloned. Several clones were sequenced and found to encode a pig *mpl* ligand of 332 amino acid resudues referred to as pML (or pML332) having the nucleotide and deduced amino acid sequence shown in Fig. 20 (SEQ ID NOS: 18 & 19).

Again, a second form, designated pML2, encoding a protein with a 4 amino acid residue deletion (228 amino acid residues) was identified (see Fig. 21 [SEQ ID NO: 21]). Comparison of pML and pML2 amino acid sequences shows the latter form is identical except that the tetrapeptide QLPP corresponding to residues 111-114 inclusive have been deleted (see Fig. 22 [SEQ ID NOS: 18 & 21]). The four amino acid deletions observed in both murine and porcine ML cDNA occur at precisely the same position within the predicted proteins.

Comparison of the predicted amino acid sequences of the mature ML from human, mouse, and pig (Fig. 19 [SEQ ID NOS: 6, 17 & 18]) indicates that overall

sequence identity is 72 percent between mouse and human, 68 percent between mouse and pig and 73 percent between pig and human. The homology is substantially greater in the amino-terminal half of the ML (EPO homologous domain). This domain is 80 to 84 percent identical between any two species whereas the carboxy-terminal half (carbohydrate domain) is only 57 to 67 percent identical. A di-basic amino acid motif that could represent a protease cleavage site is present at the carboxyl end of the erythropoeltin homology domain. This motif is conserved between the three species at this position (Fig. 19 [SEQ ID NOS: 6, 17 & 18]). A second di-basic site present at position 245 and 246 in the human sequence is not present in the mouse or pig sequences. The murine and the pig ML sequence contain 4 cysteines, all conserved in the human sequence. There are seven potential N-glycosylation sites within the mouse ligand and six within the porcine ML, 5 of which are conserved within the human sequence. Again, all the potential N-glycosylation sites are located in the C-terminal half of the protein.

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# 12. Expression and Purification of TPO from Chinese Hamster Ovary (CHO) Cells

The expression vectors used to transfect CHO cells are designated: pSVI5.ID.LL.MLORF (full length or TPO332), and pSVI5.ID.LL.MLEPO-D (truncated or TPO153). The pertinent features of these plasmids are presented in Fig. 23 and 24.

The transfection procedures are described in **Example 20**. Briefly, cDNA corresponding to the entire open reading frame of TPO was obtained by PCR. The PCR product was purified and cloned between two restriction sites (Clal and Sall) of the plasmid pSVI5.ID.LL to obtain the vector pSVI5.ID.LL.MLORF. A second construct corresponding to the EPO homologous domain was generated the same way but using a different reverse primer(EPOD.Sal). The final construct for the vector coding for the EPO homologous domain of TPO is called pSVI5.ID.LL.MLEPO-D.

These two constructs were linearized with Noti and transfected into Chinese Hamster Ovary Cells (CHO-DP12 cells, EP 307,247 published 15 March 1989) by electroporation. 10<sup>7</sup> cells were electroporated in a BRL electroporation apparatus (350 Volts, 330 mF, low capacitance) in the presence of 10, 25 or 50 mg of DNA as described (Andreason, G.L. *J. Tissue Cult. Meth.* 15,56 [1993]). The day following transfection, cells were split in DHFR selective media (High glucose DMEM-F12 50:50 without glycine, 2mM glutamine, 2-5% dialyzed fetal calf serum). 10 to 15 days later individual colonies were transferred to 96 well plates and allowed to grow to confluency. Expression of ML<sub>153</sub> or ML<sub>332</sub> in the conditioned media from these clones was assessed using the Ba/F3-mpl proliferation assay (described in Example I).

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The process for purifying and isolating TPO from harvested CHO cell-culture fluid is described in **Example 20**. Briefly, harvested cell culture fluid (HCCF) is applied to a Blue Sepharose column (Phamacia) at a ratio of approximately 100L of HCCF per liter of resin. The column is then washed with 3 to 5 column volumes of buffer followed by 3 to 5 column volumes of a buffer containing 2.0M urea. TPO is then eluted with 3 to 5 column volumes of buffer containing both 2.0M urea and 1.0M NaCl.

The Blue Sepharose eluate pool containing TPO is then applied to a Wheat Germ Lectin Sepharose column (Pharmacia) equilibrated in the Blue Sepharose eluting buffer at a ratio of from 8 to 16 ml of Blue Sepharose eluate per ml of resin. The column is then washed with 2 to 3 column volumes of equilibration buffer. TPO is then eluted with 2 to 5 column volumes of a buffer containing 2.0M urea and 0.5M N-acetyl-D-glucosamine.

The Wheat Germ Lectin eluate containing TPO is then acidified and C12E8 is added to a final concentration of 0.04%. The resulting pool is applied to a C4 reversed phase column equilibrated in 0.1% TFA, 0.04% C12E8 at a load of approximately 0.2 to 0.5 mg protein per ml of resin.

The protein is eluted in a two phase linear gradient of acetonitrile containing 0.1% TFA and 0.04% C12E8 and a pool is made on the basis of SDS-PAGE.

The C4 Pool is then diluted and diafilitered versus approximately 6 volumes of buffer on an Amicon YM or like ultrafiltration membrane having a 10,000 to 30,000 Dalton molecular weight cut-off. The resulting diafiltrate may be then directly processed or further concentrated by ultrafiltration. The diafiltrate/concentrate is usually adjusted to a final concentration of 0.01% Tween-80.

All or a portion of the diafiltrate/concentrate equivalent to 2 to 5% of the calculated column volume is then applied to a Sephacryl S-300 HR column (Pharmacia) equilibrated in a buffer containing 0.01% Tween-80 and chromatographed. The TPO containing fractions which are free of aggregate and proteolytic degradation products are then pooled on the basis of SDS-PAGE. The resulting pool is filtered and stored at 2-8°C.

# 13. Methods for Transforming and Inducing TPO Synthesis in a Microorganism and Isolating, Purifying and Refolding TPO Made Therein

Construction of *E. coli* TPO expression vectors is described in detail in **Example 21**. Briefly. plasmids pMP21, pMP151, pMP41, pMP57 and pMP202 were all designed to express the first 155 amino acids of TPO downstream of a small leader which varies among the different constructs. The leaders provide primarily for

high level translation initiation and rapid purification. The plasmids pMP210-1, -T8, -21, -22, -24, -25 are designed to express the first 153 amino acids of TPO downstream of an initiation methionine and differ only in the codon usage for the first 6 amino acids of TPO, while the plasmid pMP251 is a derivative of pMP210-1 in which the carboxy-terminal end of TPO is extended by two amino acids. All of the above plasmids will produce high levels of intracellular expression of TPO in *E. coli* upon induction of the tryptophan promoter (Yansura, D. G. et. al. Methods in Enzymology (Goeddel, D. V., Ed.) 185:54-60, Academic Press, San Diego [1990]). The plasmids pMP1 and pMP172 are intermediates in the construction of the above TPO intracellular expression plasmids.

The above TPO expression plasmids were used to transform the *E. coli* using the CaCl<sub>2</sub> heat shock method (Mandel, M. *et al. J. Mol. Biol.*, **53**:159-162, [1970]) and other procedures described in **Example 21**. Briefly, the transformed cells were grown first at 37°C until the optical density (600nm) of the culture reached approximately 2-3. The culture was then diluted and, after growth with aeration, acid was added. The culture was then allowed to continue growing with aeration for another 15 hours after which time the cells were harvested by centrifugation.

The Isolation, Purification and Refolding procedures given below for production of biologically active, refolded human TPO or fragments thereof is described in **Examples 22** and **23** can be applied for the recovery of any TPO variant including N and C terminal extended forms. Other procedures sultable for refolding recombinant or synthetic TPO can be found in the following patents; Builder *et al.*, U.S. Patent 4,511,502; Jones *et al.*, U.S. Patent 4,512,922; Olson U.S. Patent 4,518,526 and Builder *et al.*, U.S. Patent 4,620,948; for a general description of the recovery and refolding process for a variety of recombinant proteins expressed in an insoluble form in *E. coli*.

### A Recovery of non-soluble TPO

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A microorganism such as *E. coli* expressing TPO encoded by any suitable plasmid is fermented under conditions in which TPO is deposited in insoluble "refractile bodies". Optionally, cells are first washed in a cell disruption buffer. Typically, about 100g of cells are resuspended in about 10 volumes of a cell disruption buffer (e.g. 10 mM Tris, 5 mM EDTA, pH 8) with, for example, a Polytron homogenizer and the cells centrifuged at 5000 x g for 30 minutes. Cells are then lysed using any conventional technique such as tonic shock, sonication, pressure cycling, chemical or enzymatic methods. For example, the washed cell pellet above may be resuspended in another 10 volumes of a cell disruption buffer with a homogenizer and the cell suspension is passed through an LH Cell Disrupter (LH Inceltech, Inc.) or through a Microfluidizer (Microfluidics International) according to the manufactures'

instructions. The particulate matter containing TPO is then separated from the liquid phase and optionally washed with any suitable liquid. For example, a suspension of cell lysate may be centrifuged at 5,000 X g for 30 minutes, resuspended and optionally centrifuged a second time to make a washed refractile body pellet. The washed pellet may be used immediately or optionally stored frozen (at e.g. -70°C).

## B. Solubilization and Purification of Monomeric TPO

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Insoluble TPO in the refractile body pellet is then solubilized with a solublizing buffer. The solublizing buffer contains a chaotropic agent and is usually buffered at a basic pH and contains a reducing agent to improve the yield of monomeric TPO. Representative chaotropic agents include urea, guanidine HCl, and sodium thiocyanate. A preferred chaotropic agent is guanidine HCI. The concentration of chaotropic agent is usually 4-9M, preferably 6-8M. The pH of the solublizing buffer is maintained by any suitable buffer in a pH range of from about 7.5-9.5, preferably 8.0-9.0 and most preferably 8.0. Preferably the solubilizing buffer also contains a reducing agent to aid formation of the monomeric form of TPO. Suitable reducing agents include organic compounds containing a free thiol (RSH). Representative reducing agents include dithiothreitol (DTT), dithioerythritol (DTE), mercaptoethanol, glutathione (GSH), cysteamine and cysteine. A preferred reducing agent is dithiothreitol (DTT). Optionally, the solubilizing buffer may contain a mild oxidizing agent (e.g. molecular oxygen) and a sulfite salt to form monomeric TPO via sulfitolysis. In this embodiment, the resulting TPO-S-sulfonate is later refolded in the presence of the redox buffer (e.g. GSH/GSSG) to form the properly folded TPO.

The TPO protein is usually further purified using, for example, centrifugation, gel filtration chromatography and reversed phase column chromatography.

By way of illustration, the following procedure has produced suitable yields of monomeric TPO. The refractile body pellet is resuspended in about 5 volumes by weight of the solubilizing buffer (20 mM Tris, pH 8, with 6-8 M guanidine and 25 mM DTT) and stirred for 1-3 hr., or overnight, at 4°C to effect solubilization of the TPO protein. High concentrations of urea (6-8M) are also useful but generally result in somewhat lower yields compared to guanidine. After solubilization, the solution is centrifuged at 30,000 x g for 30 min. to produce a clear supernatant containing denatured, monomeric TPO protein. The supernatant is then chromatographed on a Superdex 200 gel filtration column (Pharmacia, 2.6 x 60 cm) at a flow rate of 2 ml/min. and the protein eluted with 20 mM Na phosphate, pH 6.0, with 10 mM DTT. Fractions containing monomeric, denatured TPO protein eluting between 160 and 200 ml are pooled. The TPO protein is further purified on a semi-preparative C4 reversed phase column (2 x 20 cm VYDAC). The sample is applied at 5 ml/min. to a column equilibrated in 0.1% TFA (trifluoroacetic acid) with 30% acetonitrile. The protein is

eluted with a linear gradient of acetonitrile (30-60% in 60 min.). The purified reduced protein elutes at approximately 50% acetonitrile. This material is used for refolding to obtain biologically active TPO variant.

### C Refolding TPO to Generate the Biologically Active Form

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Following solubilization and further purification of TPO, the biologically active form is obtained by refolding the denatured monomeric TPO in a redox buffer. Because of the high potency of TPO (half maximal stimulation in the Ba/F3 assay is achieved at approximately 3 pg/ml), it is possible to obtain biologically active material utilizing many different buffer, detergent and redox conditions. However, under most conditions only a small amount of properly folded material (<10%) is obtained. For commercial manufacturing processes, it is desirable to have refolding yields at least 10%, more preferably 30-50% and most preferably >50%. Many different detergents including Triton X-100, dodecyl-beta-maltoside, CHAPS, CHAPSO, SDS, sarkosyl, Tween 20 and Tween 80, Zwittergent 3-14 and others were found suitable for producing at least some properly folded material. Of these however, the most preferred detergents were those of the CHAPS family (CHAPS and CHAPSO) which were found to work best in the refolding reaction and to limit protein aggregation and improper disulfide formation. Levels of CHAPS greater than about 1% were most preferred. Sodium chloride was required for the best yields, with the optimal levels between 0.1 M and 0.5M. The presence of EDTA (1-5 mM) in the redox buffer was preferred to limit the amount of metal-catalyzed oxidation (and aggregation) which was observed with some preparations. Glycerol concentrations of greater than 15% produced the optimal refolding conditions. For maximum ylelds, it was essential to have a redox pair in the redox buffer consisting of both an oxidized and reduced organic thiol (RSH). Suitable redox pairs include mercaptoethanol, glutathione (GSH), cysteamine, cysteine and their corresponding oxidized forms. Preferred redox pairs were glutathione(GSH):oxidized glutathione(GSSG) or cysteine:cystine. The most preferred redox pair was glutathione(GSH):oxidized glutathione(GSSG). Generally higher yields were observed when the mole ratio of oxidized member of the redox pair was equal to or in excess over the reduced member of the redox pair. pH values between 7.5 and about 9 were optimal for refolding of these TPO variants. Organic solvents (e.g. ethanol, acetonitrile, methanol) were tolerated at concentrations of 10-15% or lower. Higher levels of organic solvents increased the amount of improperly folded forms. Tris and phosphate buffers were generally useful. Incubation at 4 °C also produced higher levels of properly folded TPO.

Refolding yields of 40-60% (based on the amount of reduced and denatured TPO used in the refolding reaction) are typical for preparations of TPO that have been purified through the first C4 step. Active material can be obtained when less pure

preparations (e.g. directly after the Superdex 200 column or after the initial refractile body extraction) although the yields are less due to extensive precipitation and interference of non-TPO proteins during the TPO refolding process.

Since TPO contains 4 cysteine residues, it is possible to generate three different disulfide versions of this protein:

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version 1: disulfides between cysteine residues 1-4 and 2-3

version 2: disulfides between cysteine residues 1-2 and 3-4

version 3: disulfides between cysteine residues 1-3 and 2-4.

During the initial exploration in determining refolding conditions, several different peaks containing the TPO protein were separated by C4 reversed phase chromatography. Only one of these peaks had significant biological activity as determined using the Ba/F3 assay. Subsequently, the refolding conditions were optimized to yield preferentially that version. Under these conditions, the misfolded versions were less than 10-20% of the total monomeric TPO obtained from the solubilizing step.

The disulfide pattern for the biologically active TPO has been determined to be 1-4 and 2-3 by mass spectrometry and protein sequencing, where the cysteines are numbered sequentially from the amino-terminus. This cysteine cross-linking pattern is consistent with the known disulfide bonding pattern of the related molecule erythropoietin.

# D. Biological Activity of Recombinant, Refolded TPO

Refolded and purified TPO has activity in both *in vitro* and *in vivo* assays. For example, in the Ba/F3 assay, half-maximal stimulation of thymidine incorporation into the Ba/F3 cells for TPO (Met<sup>-1</sup> 1-153) was achieved at 3.3 pg /ml (0.3 pM). In the *mpl* receptor-based ELISA, half-maximal activity occurred at 1.9 ng/ml (120 pM). In normal and myelosuppressed animals produced by near-lethal X-radiation, refolded TPO (Met<sup>-1</sup> 1-153) was highly potent (activity was seen at doses as low as 30 ng/mouse) to stimulate the production of new platelets. Similar biological activity was observed for other forms of TPO refolded in accordance with the above described procedures (see Figs. 25, 26 and 28).

# 14. Methods for Measurement of Thrombopoletic Activity

Thrombopoietic activity may be measured in various assays including the Ba/F3 mpl ligand assay described in Example 1, an in vivo mouse platelet rebound synthesis assay, induction of platelet cell surface antigen assay as measured by an anti-platelet immunoassay (anti-GPIIbIIIa) for a human leukemia megakaryoblastic cell line (CMK) (see Sato et al., Brit. J. Heamatol., 72:184-190 [1989])(see also the liquid suspension megakaryocytopoiesis assay described in Example 4), and

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induction of polyploidization in a megakaryoblastic cell line (DAMI) (see Ogura et al., Blood, 72(1):49-60 [1988]). Maturation of megakaryocytes from immature, largely non-DNA synthesizing cells, to morphologically identifiable megakaryocytes involves a process that includes appearance of cytoplasmic organelles, acquisition of membrane antigens (GPIIbIIIa), endoreplication and release of platelets as described in the background. A lineage specific promoter (i.e., the mpl ligand) of megakaryocyte maturation would be expected to induce at least some of these changes in immature megakaryocytes leading to platelet release and alleviation of thrombocytopenia. Thus, assays were designed to measure the emergence of these parameters in immature megakaryocyte cell lines, i.e., CMK and DAMI cells. The CMK assay (Example 4) measures the appearance of a specific platelet marker, GPIIbIIIa, and platelet shedding. The DAMI assay (Example 15) measures endoreplication since increases in ploidy are hallmarks of mature megakaryocytes. Recognizable megakaryocytes have ploidy values of 2N, 4N, 8N, 16N, 32N, etc. Finally, the in vivo mouse platelet rebound assay (Example 16) is useful in demonstrating that administration of the test compound (here the mpl ligand) results in elevation of platelet numbers.

Two additional *in vitro* assays have been developed to measure TPO activity. The first is a kinase receptor activation (KIRA) ELISA in which CHO cells are transfected with a *mpl*-Rse chimera and tyrosine phosphorylation of Rse is measured by ELISA after exposure of the *mpl* portion of the chimera to *mpl* ligand (see Example 17). The second is a receptor based ELISA in which ELISA plate coated rabbit antihuman IgG captures human chimeric receptor *mpl*-IgG which binds the *mpl* ligand being assayed. A biotinylated rabbit polyclonal antibody to *mpl* ligand (TPO<sub>155</sub>) is used to detect bound *mpl* ligand which is measured using streptavidin-peroxidase as described in Example 18.

# 15. In Vivo Biological Response of Normal and Sublethally Irradiated Mice Treated with TPO

Both normal and sublethally irradiated mice were treated with truncated and full length TPO isolated from Chinese hamster ovary (CHO) cells, *E. coli*, and human embryonic kidney (293) cells. Both forms of TPO produced in these three hosts stimulated platelet production in mice, however, full length TPO isolated from CHO apeared to produce the greatest *in vivo* response. These results indicate that proper glycosylation of the carboxy-terminal domain may be necessary for optimal *in vivo* activity.

## (a) E. coli-rhTPO(Met<sup>-1</sup>,153)

The "Met" form of the EPO domain (Met in the -1 position plus the first 153 residues of human TPO) produced in *E. coli* (see Example 23) was injected daily into

normal female C57 B6 mice as described in the legends to Figs. 25A, 25B and 25C. These figures show that the non-glycosylated truncated form of TPO produced in *E. coli* and refolded as described above is capable of stimulating about a two-fold increase in platelet production in normal mice with out effecting the red or white blood cell population.

This same molecule injected daily into sublethally irradiated (137Cs) female C57 B6 mice as described in the legends to Figs. 26A, 26B and 26C stimulated platelet recovery and diminished nadir but had no effect on erythrocytes or leukocytes.

### (b) CHO-rhTPO332

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The full length form of TPO produced in CHO and injected daily into normal female C57 B6 mice as described in the legends to Figs. 27A, 27B and 27C produced about a five-fold increase in platelet production in normal mice with out effecting the erythrocyte or leukocyte population.

(c) CHO-rhTPO332; E. coli-rhTPO(Met<sup>-1</sup>,153); 293-rhTPO332; and E. coli-rhTPO155

Dose response curves were constructed for treatment of normal mice with rhTPO from various cell lines (CHO-rhTPO332; *E. coli*-rhTPO(Met<sup>-1</sup>,153); 293-rhTPO332; and *E. coli*-rhTPO155) as described in the legend to **Fig. 28**. This figure shows that all tested forms of the molecule stimulate platelet production, however the full length form produced in CHO has the greatest *in vivo* activity.

## (d) CHO-rhTPO<sub>153</sub>, CHO-rhTPO<sub>"clipped"</sub> and CHO-rhTPO<sub>332</sub>

Dose response curves were also constructed for treatment of normal mice with various forms of rhTPO produced in CHO (CHO-rhTPO153, CHO-rhTPO"clipped" and CHO-rhTPO332) as described in the legend to Fig. 29. This figure shows that all tested CHO forms of the molecule stimulate platelet production, but that the full length 70 Kda form has the greatest *in vivo* activity.

# 16. General Recombinant Preparation of *mpl* Ligand and Variants

Preferably mpl ligand is prepared by standard recombinant procedures which involve production of the mpl ligand polypeptide by culturing cells transfected to express mpl ligand nucleic acid (typically by transforming the cells with an expression vector) and recovering the polypeptide from the cells. However, it is optionally envisioned that the mpl ligand may be produced by homologous recombination, or with recombinant production methods utilizing control elements introduced into cells already containing DNA encoding the mpl ligand. For example, a powerful promoter/enhancer element, a suppressor, or an exogenous transcription modulatory element may be inserted in the genome of the intended host cell in

proximity and orientation sufficient to influence the transcription of DNA encoding the desired *mpl* ligand polypeptide. The control element does not encode the *mpl* ligand, rather the DNA is indigenous to the host cell genome. One next screens for cells making the receptor polypeptide of this invention, or for increased or decreased levels of expression, as desired.

Thus, the invention contemplates a method for producing *mpl* ligand comprising inserting into the genome of a cell containing the *mpl* ligand nucleic acid molecule a transcription modulatory element in sufficient proximity and orientation to the nucleic acid molecule to influence transcription thereof, with an optional further step comprising culturing the cell containing the transcription modulatory element and the nucleic acid molecule. The invention also contemplates a host cell containing the indigenous *mpl* ligand nucleic acid molecule operably linked to exogenous control sequences recognized by the host cell.

## A. Isolation of DNA Encoding mpl ligand Polypeptide

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The DNA encoding *mpl* ligand polypeptide may be obtained from any cDNA library prepared from tissue believed to possess the *mpl* ligand mRNA and to express it at a detectable level. The *mpl* ligand gene may also be obtained from a genomic DNA library or by *in vitro* oligonucleotide synthesis from the complete nucleotide or amino acid sequence.

Libraries are screened with probes designed to identify the gene of interest or the protein encoded by it. For cDNA expression libraries, suitable probes include monoclonal or polyclonal antibodies that recognize and specifically bind to the *mpl* ligand. For cDNA libraries suitable probes include oligonucleotides of about 20-80 bases in length that encode known or suspected portions of the *mpl* ligand cDNA from the same or different species; and/or complementary or homologous cDNAs or fragments thereof that encode the same or a similar gene. Appropriate probes for screening genomic DNA libraries include, but are not limited to, oligonucleotides, cDNAs, or fragments thereof that encode the same or a similar gene, and/or homologous genomic DNAs or fragments thereof. Screening the cDNA or genomic library with the selected probe may be conducted using standard procedures as described in Chapters 10-12 of Sambrook *et al.*, *supra*.

An alternative means to isolate the gene encoding *mpl* ligand is to use PCR methodology as described in section 14 of Sambrook *et al.*, *supra*. This method requires the use of oligonucleotide probes that will hybridize to DNA encoding the *mpl* ligand. Strategies for selection of oligonucleotides are described below.

A preferred method of practicing this invention is to use carefully selected oligonucleotide sequences to screen cDNA libraries from various tissues, preferably human or porcine kidney (adult or fetal) or liver cell lines. For example, human fetal

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liver cell line cDNA libraries are screened with the oligonucleotide probes. Alternatively, human genomic libraries may be screened with the oligonucleotide probes.

The oligonucleotide sequences selected as probes should be of sufficient length and sufficiently unambiguous that false positives are minimized. The actual nucleotide sequence(s) is usually designed based on regions of the *mpl* ligand which have the least codon redundancy. The oligonucleotides may be degenerate at one or more positions. The use of degenerate oligonucleotides is of particular importance where a library is screened from a species in which preferential codon usage is not known.

The oligonucleotide must be labeled such that it can be detected upon hybridization to DNA in the library being screened. The preferred method of labeling is to use ATP  $(e.g., \gamma^{32}P)$  and polynucleotide kinase to radiolabel the 5' end of the oligonucleotide. However, other methods may be used to label the oligonucleotide, including, but not limited to, biotinylation or enzyme labeling.

Of particular interest is the *mpl* ligand nucleic acid that encodes a full-length *mpl* ligand polypeptide. In some preferred embodiments, the nucleic acid sequence includes the native *mpl* ligand signal sequence. Nucleic acid having all the protein coding sequence is obtained by screening selected cDNA or genomic libraries using the deduced amino acid sequence.

## B. Amino Acid Sequence Variants of Native mpl ligand

Amino acid sequence variants of mpl ligand are prepared by introducing appropriate nucleotide changes into the mpl ligand DNA, or by in vitro synthesis of the desired mpl ligand polypeptide. Such variants include, for example, deletions from, or insertions or substitutions of, residues within the amino acid sequence for the porcine mpl ligand. For example, carboxy terminus portions of the mature full length mpl ligand may be removed by proteolytic cleavage, either in vivo or in vitro, or by cloning and expressing a fragment or the DNA encoding full length mpl ligand to produce a biologically active variant. Any combination of deletion, insertion, and substitution is made to arrive at the final construct, provided that the final construct possesses the desired biological activity. The amino acid changes also may alter posttranslational processes of the mpl ligand, such as changing the number or position of glycosylation sites. For the design of amino acid sequence variants of the mpl ligand, the location of the mutation site and the nature of the mutation will depend on the mpl ligand characteristic(s) to be modified. The sites for mutation can be modified individually or in series, e.g., by (1) substituting first with conservative amino acid choices and then with more radical selections depending upon the results achieved, (2) deleting the target residue, or (3) inserting residues of the same or a different class adjacent to the located site, or combinations of options 1-3.

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A useful method for identification of certain residues or regions of the *mpl* ligand polypeptide that are preferred locations for mutagenesis is called "alanine scanning mutagenesis," as described by Cunningham and Wells, *Science*, 244:1081-1085 [1989]. Here, a residue or group of target residues are Identified (*e.g.*, charged residues such as arg, asp, his, lys, and glu) and replaced by any, but preferably a neutral or negatively charged, amino acid (most preferably alanine or polyalanine) to affect the interaction of the amino acids with the surrounding aqueous environment in or outside the cell. Those domains demonstrating functional sensitivity to the substitutions then are refined by introducing further or other variants at or for the sites of substitution. Thus, while the site for introducing an amino acid sequence variation is predetermined, the nature of the mutation *per se* need not be predetermined. For example, to optimize the performance of a mutation at a given site, ala scanning or random mutagenesis is conducted at the target codon or region and the expressed *mpl* ligand variants are screened for the optimal combination of desired activity.

There are two principal variables in the construction of amino acid sequence variants: the location of the mutation site and the nature of the mutation. For example, variants of the *mpl* ligand polypeptide include variants from the *mpl* ligand sequence, and may represent naturally occurring alleles (which will not require manipulation of the *mpl* ligand DNA) or predetermined mutant forms made by mutating the DNA, either to arrive at an allele or a variant not found in nature. In general, the location and nature of the mutation chosen will depend upon the *mpl* ligand characteristic to be modified.

Amino acid sequence deletions generally range from about 1 to 30 residues, more preferably about 1 to 10 residues, and typically are contiguous. Alternatively, amino acid sequence deletions for the *mpl* ligand may include a portion of or the entire carboxy-terminus glycoprotein domain. Amino acid sequence deletions may also include one or more of the first 6 amino-terminus residues of the mature protein. Optional amino acid sequence deletions comprise one or more residues in one or more of the loop regions that exist between the 'helical bundels''. Contiguous deletions ordinarily are made in even numbers of residues, but single or odd numbers of deletions are within the scope hereof. Deletions may be introduced into regions of low homology among the *mpl* ligands that share the most sequence identity to modify the activity of the *mpl* ligand. Or deletions may be introduced into regions of low homology among human *mpl* ligand and other mammalian *mpl* ligand polypeptides that share the most sequence identity to the human *mpl* ligand. Deletions from a mammalian *mpl* ligand polypeptide in areas of substantial homology with other mammalian *mpl* ligands will be more likely to modify the biological activity of the *mpl* ligand more

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significantly. The number of consecutive deletions will be selected so as to preserve the tertiary structure of *mpl* ligands in the affected domain, *e.g.*, beta-pleated sheet or alpha helix.

Amino acid sequence insertions include amino- and/or carboxyl-terminal fusions ranging in length from one residue to polypeptides containing a hundred or more residues, as well as intrasequence insertions of single or multiple amino acid residues. Intrasequence insertions (i.e., insertions within the mature mpl ligand sequence) may range generally from about 1 to 10 residues, more preferably 1 to 5, most preferably 1 to 3. An exemplary preferred fusion is that of mpl ligand or fragment thereof and another cytokine or fragment thereof. Examples of terminal insertions include mature mpl ligand with an N-terminal methionyl residue, an artifact of the direct expression of mature mpl ligand in recombinant cell culture, and fusion of a heterologous N-terminal signal sequence to the N-terminus of the mature mpl ligand molecule to facilitate the secretion of mature mpl ligand from recombinant hosts. Such signal sequences generally will be obtained from, and thus homologous to, the intended host cell species. Suitable sequences include STII or lpp for E. coli, alpha factor for yeast, and viral signals such as herpes gD for mammalian cells.

Other insertional variants of the *mpl* ligand molecule include the fusion to the N- or C-terminus of *mpl* ligand of immunogenic polypeptides (*i.e.*, not endogenous to the host to which the fusion is administered), *e.g.*, bacterial polypeptides such as beta-lactamase or an enzyme encoded by the *E. coli trp* locus, or yeast protein, and C-terminal fusions with proteins having a long half-life such as immunoglobulin constant regions (or other immunoglobulin regions), albumin, or ferritin, as described in WO 89/02922 published 6 April 1989.

A third group of variants are amino acid substitution variants. These variants have at least one amino acid residue in the *mpl* ligand molecule removed and a different residue inserted in its place. The sites of greatest interest for substitutional mutagenesis include sites identified as the active site(s) of *mpl* ligand and sites where the amino acids found in other analogues are substantially different in terms of sidechain bulk, charge, or hydrophobicity, but where there is also a high degree of sequence identity at the selected site among various *mpl* ligand species and/or within the various animal analogues of one *mpl* ligand member.

Other sites of interest are those in which particular residues of the *mpl* ligand obtained from various family members and/or animal species within one member are identical. These sites, especially those falling within a sequence of at least three other identically conserved sites, are substituted in a relatively conservative manner. Such conservative substitutions are shown in Table 3 under the heading of preferred substitutions. If such substitutions result in a change in biological activity, then more

substantial changes, denominated exemplary substitutions in **Table 3**, or as further described below in reference to amino acid classes, are introduced and the products screened.

5	TABLE 3						
	Original	Exemplary	Preferred				
	Residue	Substitutions	Substitutions				
	Ala (A)	Val; Leu; Ile	Val				
	Arg (R)	Lys; Gln; Asn	Lys				
10	Asn (N)	Gin; His; Lys; Arg	Gln				
	Asp (D)	Glu	Glu				
	Cys (C)	Ser	Ser				
	Gin (Q)	Asn	Asn				
	Glu (E)	Asp	Asp				
15	Gly (G)	Pro	Pro				
	His (H)	Asn; Gln; Lys; Arg	Arg				
	lle (I)	Leu; Val; Met; Ala; Phe	<b>;</b> ;				
		norleucine	Leu				
	Leu (L)	norleucine; Ile; Val;					
20		Met; Ala; Phe	lle				
	Lys (K)	Arg; Gln; Asn	Arg				
	Met (M)	Leu; Phe; Ile	Leu				
	Phe (F)	Leu; Val; Ile; Ala	Leu				
	Pro (P)	Gly	Gly				
25	Ser (S)	Thr	Thr				
	Thr (T)	Ser	Ser				
	Trp (W)	Tyr	Tyr				
	Tyr (Y)	Trp; Phe; Thr; Ser	Phe				
	Val (V)	lle; Leu; Met; Phe;					
30		Ala; norleucine	Leu				

Substantial modifications in function or immunological identity of the *mpl* ligand are accomplished by selecting substitutions that differ significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Naturally occurring residues are divided into groups based on common side-chain properties:

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(1) hydrophobic: norleucine, Met, Ala, Val, Leu, Ile;

(2) neutral hydrophilic: Cys, Ser, Thr;

(3) acidic: Asp, Glu;

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(4) basic: Asn, Gln, His, Lys, Arg;

(5) residues that influence chain orientation: Gly, Pro; and

(6) aromatic: Trp, Tyr, Phe.

Non-conservative substitutions will entail exchanging a member of one of these classes for another. Such substituted residues also may be introduced into the conservative substitution sites or, more preferably, into the remaining (non-conserved) sites.

In one embodiment of the invention, it is desirable to inactivate one or more protease cleavage sites that are present in the molecule. These sites are identified by inspection of the encoded amino acid sequence, in the case of trypsin, e.g., for an arginyl or lysinyl residue. When protease cleavage sites are identified, they are rendered inactive to proteolytic cleavage by substituting the targeted residue with another residue, preferably a basic residue such as glutamine or a hydrophobic residue such as serine; by deleting the residue; or by inserting a prolyl residue immediately after the residue.

In another embodiment, any methionyl residues other than the starting methionyl residue of the signal sequence, or any residue located within about three residues N- or C-terminal to each such methionyl residue, is substituted by another residue (preferably in accordance with **Table 3**) or deleted. Alternatively, about 1-3 residues are inserted adjacent to such sites.

Any cysteine residues not involved in maintaining the proper conformation of the *mpl* ligand also may be substituted, generally with serine, to improve the oxidative stability of the molecule and prevent aberrant crosslinking. It has been found that the first and forth cysteines in the epo domain, numbered from the amino-terminus, are necessary for maintaining proper conformation but that the second and third are not. Accordingly, the second and third cysteines in the epo domain may be substituted.

Nucleic acid molecules encoding amino acid sequence variants of *mpl* ligand are prepared by a variety of methods known in the art. These methods include, but are not limited to, isolation from a natural source (in the case of naturally occurring amino acid sequence variants) or preparation by oligonucleotide-mediated (or site-directed) mutagenesis, PCR mutagenesis, and cassette mutagenesis of an earlier prepared variant or a non-variant version of *mpl* ligand polypeptide.

Oligonucleotide-mediated mutagenesis is a preferred method for preparing substitution, deletion, and insertion variants of *mpl* ligand DNA. This technique is well known in the art as described by Adelman *et al.*, *DNA*, 2:183 [1983]. Briefly, *mpl* 

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ligand DNA is altered by hybridizing an oligonucleotide encoding the desired mutation to a DNA template, where the template is the single-stranded form of a plasmid or bacteriophage containing the unaltered or native DNA sequence of *mpl* ligand. After hybridization, a DNA polymerase is used to synthesize an entire second complementary strand of the template that will thus incorporate the oligonucleotide primer, and will code for the selected alteration in the *mpl* ligand DNA.

Generally, oligonucleotides of at least 25 nucleotides in length are used. An optimal oligonucleotide will have 12 to 15 nucleotides that are completely complementary to the template on either side of the nucleotide(s) coding for the mutation. This ensures that the oligonucleotide will hybridize properly to the single-stranded DNA template molecule. The oligonucleotides are readily synthesized using techniques known in the art such as that described by Crea et al., Proc. Natl. Acad. Sci. USA, 75:5765 [1978].

The DNA template can be generated by those vectors that are either derived from bacteriophage M13 vectors (the commercially available M13mp18 and M13mp19 vectors are suitable), or those vectors that contain a single-stranded phage origin of replication as described by Viera et al., Meth. Enzymol., 153:3 [1987]. Thus, the DNA that is to be mutated may be inserted into one of these vectors to generate single-stranded template. Production of the single-stranded template is described in Sections 4.21-4.41 of Sambrook et al., Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Laboratory Press, NY 1989).

Alternatively, single-stranded DNA template may be generated by denaturing double-stranded plasmid (or other) DNA using standard techniques.

For alteration of the native DNA sequence (to generate amino acid sequence variants, for example), the oligonucleotide is hybridized to the single-stranded template under suitable hybridization conditions. A DNA polymerizing enzyme, usually the Klenow fragment of DNA polymerase I, is then added to synthesize the complementary strand of the template using the oligonucleotide as a primer for synthesis. A heteroduplex molecule is thus formed such that one strand of DNA encodes the mutated form of the *mpl* ligand, and the other strand (the original template) encodes the native, unaltered sequence of the *mpl* ligand. This heteroduplex molecule is then transformed into a suitable host cell, usually a prokaryote such as *E. coli* JM101. After the cells are grown, they are plated onto agarose plates and screened using the oligonucleotide primer radiolabeled with 32-phosphate to identify the bacterial colonies that contain the mutated DNA. The mutated region is then removed and placed in an appropriate vector for protein production, generally an expression vector of the type typically employed for transformation of an appropriate host.

The method described immediately above may be modified such that a homoduplex molecule is created wherein both strands of the plasmid contain the mutation(s). The modifications are as follows: The single-stranded oligonucleotide is annealed to the single-stranded template as described above. A mixture of three deoxyribonucleotides, deoxyriboadenosine (dATP), deoxyriboguanosine (dGTP), and deoxyribothymidine (dTTP), is combined with a modified thio-deoxyribocytosine called dCTP-(aS) (which can be obtained from the Amersham Corporation). This mixture is added to the template-oligonucleotide complex. Upon addition of DNA polymerase to this mixture, a strand of DNA identical to the template, except for the mutated bases is generated. In addition, this new strand of DNA will contain dCTP-(aS) instead of dCTP, which serves to protect it from restriction endonuclease digestion.

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After the template strand of the double-stranded heteroduplex is nicked with an appropriate restriction enzyme, the template strand can be digested with *ExoIII* nuclease or another appropriate nuclease past the region that contains the site(s) to be mutagenized. The reaction is then stopped to leave a molecule that is only partially single-stranded. A complete double-stranded DNA homoduplex is then formed using DNA polymerase in the presence of all four deoxyribonucleotide triphosphates, ATP, and DNA ligase. This homoduplex molecule can then be transformed into a suitable host cell such as *E. coli* JM101, as described above.

DNA encoding *mp1* ligand mutants with more than one amino acid to be substituted may be generated in one of several ways. If the amino acids are located close together in the polypeptide chain, they may be mutated simultaneously using one oligonucleotide that codes for all of the desired amino acid substitutions. If, however, the amino acids are located some distance from each other (separated by more than about ten amino acids), it is more difficult to generate a single oligonucleotide that encodes all of the desired changes. Instead, one of two alternative methods may be employed.

In the first method, a separate oligonucleotide is generated for each amino acid to be substituted. The oligonucleotides are then annealed to the single-stranded template DNA simultaneously, and the second strand of DNA that is synthesized from the template will encode all of the desired amino acid substitutions.

The alternative method involves two or more rounds of mutagenesis to produce the desired mutant. The first round is as described for the single mutants: wild-type DNA is used for the template, an oligonucleotide encoding the first desired amino acid substitution(s) is annealed to this template, and the heteroduplex DNA molecule is then generated. The second round of mutagenesis utilizes the mutated DNA produced in the first round of mutagenesis as the template. Thus, this template already contains one or more mutations. The oligonucleotide encoding the additional desired amino acid

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substitution(s) is then annealed to this template, and the resulting strand of DNA now encodes mutations from both the first and second rounds of mutagenesis. This resultant DNA can be used as a template in a third round of mutagenesis, and so on.

PCR mutagenesis is also suitable for making amino acid variants of mpl ligand polypeptide. While the following discussion refers to DNA, it is understood that the technique also finds application with RNA. The PCR technique generally refers to the following procedure (see Erlich, supra, the chapter by R. Higuchi, p. 61-70): When small amounts of template DNA are used as starting material in a PCR, primers that differ slightly in sequence from the corresponding region in a template DNA can be used to generate relatively large quantities of a specific DNA fragment that differs from the template sequence only at the positions where the primers differ from the template. For introduction of a mutation into a plasmid DNA, one of the primers is designed to overlap the position of the mutation and to contain the mutation; the sequence of the other primer must be identical to a stretch of sequence of the opposite strand of the plasmid, but this sequence can be located anywhere along the plasmid DNA. It is preferred, however, that the sequence of the second primer is located within 200 nucleotides from that of the first, such that in the end the entire amplified region of DNA bounded by the primers can be easily sequenced. PCR amplification using a primer pair like the one just described results in a population of DNA fragments that differ at the position of the mutation specified by the primer, and possibly at other positions, as template copying is somewhat error-prone.

If the ratio of template to product material is extremely low, the vast majority of product DNA fragments incorporate the desired mutation(s). This product material is used to replace the corresponding region in the plasmid that served as PCR template using standard DNA technology. Mutations at separate positions can be introduced simultaneously by either using a mutant second primer, or performing a second PCR with different mutant primers and ligating the two resulting PCR fragments simultaneously to the vector fragment in a three (or more)-part ligation.

In a specific example of PCR mutagenesis, template plasmid DNA (1 μg) is linearized by digestion with a restriction endonuclease that has a unique recognition site in the plasmid DNA outside of the region to be amplified. Of this material, 100 ng is added to a PCR mixture containing PCR buffer, which contains the four deoxynucleotide triphosphates and is included in the GeneAmp® kits (obtained from Perkin-Elmer Cetus, Norwalk, CT and Emeryville, CA), and 25 pmole of each oligonucleotide primer, to a final volume of 50 μl. The reaction mixture is overlayed with 35 μl mineral oil. The reaction mixture is denatured for five minutes at 100°C, placed briefly on ice, and then 1 μl *Thermus aquaticus (Taq)* DNA polymerase (5 units/μl, purchased from Perkin-Elmer Cetus) is added below the mineral oil layer.

The reaction mixture is then inserted into a DNA Thermal Cycler (purchased from Perkin-Elmer Cetus) programmed as follows:

2 min. 55°C

30 sec. 72°C, then 19 cycles of the following:

30 sec. 94°C

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30 sec. 55°C, and

30 sec. 72°C.

At the end of the program, the reaction vial is removed from the thermal cycler and the aqueous phase transferred to a new vial, extracted with phenol/chloroform (50:50 vol), and ethanol precipitated, and the DNA is recovered by standard procedures. This material is subsequently subjected to the appropriate treatments for insertion into a vector.

Another method for preparing variants, cassette mutagenesis, is based on the technique described by Wells et al., Gene, 34:315 [1985]. The starting material is the plasmid (or other vector) comprising the mpl ligand DNA to be mutated. The codon(s) in the mpl ligand DNA to be mutated are identified. There must be a unique restriction endonuclease site on each side of the identified mutation site(s). If no such restriction sites exist, they may be generated using the above-described oligonucleotide-mediated mutagenesis method to introduce them at appropriate locations in the mpl ligand DNA. After the restriction sites have been introduced into the plasmid, the plasmid is cut at these sites to linearize it. A double-stranded oligonucleotide encoding the sequence of the DNA between the restriction sites but containing the desired mutation(s) is synthesized using standard procedures. The two strands are synthesized separately and then hybridized together using standard techniques. This double-stranded oligonucleotide is referred to as the cassette. This cassette is designed to have 3' and 5' ends that are compatible with the ends of the linearized plasmid, such that it can be directly ligated to the plasmid. This plasmid now contains the mutated mpl ligand DNA sequence.

## C. Insertion of Nucleic Acid Into a Replicable Vector

The nucleic acid (e.g., cDNA or genomic DNA) encoding native or variant mpl ligand polypeptide is inserted into a replicable vector for further cloning (amplification of the DNA) or for expression. Many vectors are available, and selection of the appropriate vector will depend on (1) whether it is to be used for DNA amplification or for DNA expression, (2) the size of the nucleic acid to be inserted into the vector, and (3) the host cell to be transformed with the vector. Each vector contains various components depending on its function (amplification of DNA or expression of DNA) and the host cell with which it is compatible. The vector components generally include, but are not limited to, one or more of the following: a

signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence.

#### (i) Signal Sequence Component

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The mpl ligand of this invention may be expressed not only directly, but also as a fusion with a heterologous polypeptide, preferably a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. In general, the signal sequence may be a component of the vector, or it may be a part of the mpl ligand DNA that is inserted into the vector. The heterologous signal sequence selected should be one that is recognized and processed (i.e., cleaved by a signal peptidase) by the host cell. For prokaryotic host cells that do not recognize and process the native mpl ligand signal sequence, the signal sequence is substituted by a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the native signal sequence may be substituted by, e.g., the yeast invertase, alpha factor, or acid phosphatase leaders, the C. albicans glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cell expression the native signal sequence (i.e., the mpl ligand presequence that normally directs secretion of mpl ligand from its native mammalian cells in vivo) is satisfactory, although other mammalian signal sequences may be suitable, such as signal sequences from other mpl ligand polypeptides or from the same mpl ligand from a different animal species, signal sequences from a mpl ligand, and signal sequences from secreted polypeptides of the same or related species, as well as viral secretory leaders, for example, the herpes simplex gD signal.

#### (ii) Origin of Replication Component

Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Generally, in cloning vectors this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes origins of replication or autonomously replicating sequences. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2  $\mu$  plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (the SV40 origin may typically be used only because it contains the early promoter).

Most expression vectors are "shuttle" vectors, i.e., they are capable of replication in at least one class of organisms but can be transfected into another

organism for expression. For example, a vector is cloned in *E. coli* and then the same vector is transfected into yeast or mammalian cells for expression even though it is not capable of replicating independently of the host cell chromosome.

DNA may also be amplified by insertion into the host genome. This is readily accomplished using *Bacillus* species as hosts, for example, by including in the vector a DNA sequence that is complementary to a sequence found in *Bacillus* genomic DNA. Transfection of *Bacillus* with this vector results in homologous recombination with the genome and insertion of *mpl* ligand DNA. However, the recovery of genomic DNA encoding *mpl* ligand is more complex than that of an exogenously replicated vector because restriction enzyme digestion is required to excise the *mpl* ligand DNA.

## (iii) Selection Gene Component

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Expression and cloning vectors should contain a selection gene, also termed a selectable marker. This gene encodes a protein necessary for the survival or growth of transformed host cells grown in a selective culture medium. Host cells not transformed with the vector containing the selection gene will not survive in the culture medium. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g., the gene encoding D-alanine racemase for Bacilli.

One example of a selection scheme utilizes a drug to arrest growth of a host cell. Those cells that are successfully transformed with a heterologous gene express a protein conferring drug resistance and thus survive the selection regimen. Examples of such dominant selection use the drugs neomycin (Southern et al., J. Molec. Appl. Genet., 1:327 [1982]) mycophenolic acid (Mulligan et al., Science, 209:1422 [1980]) or hygromycin Sugden et al., Mol. Cell. Biol., 5:410-413 [1985]). The three examples given above employ bacterial genes under eukaryotic control to convey resistance to the appropriate drug G418 or neomycin (geneticin), xgpt (mycophenolic acid), or hygromycin, respectively.

Examples of other suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the *mpl* ligand nucleic acid, such as dihydrofolate reductase (DHFR) or thymidine kinase. The mammalian cell transformants are placed under selection pressure that only the transformants are uniquely adapted to survive by virtue of having taken up the marker. Selection pressure is imposed by culturing the transformants under conditions in which the concentration of selection agent in the medium is successively changed, thereby leading to amplification of both the selection gene and the DNA that encodes *mpl* ligand polypeptide. Amplification is the process by which genes in greater demand for the production of a protein critical for growth are reiterated in tandem within the

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chromosomes of successive generations of recombinant cells. Increased quantities of mpl ligand are synthesized from the amplified DNA.

For example, cells transformed with the DHFR selection gene are first identified by culturing all of the transformants in a culture medium that contains methotrexate (Mtx), a competitive antagonist of DHFR. An appropriate host cell when wild-type DHFR is employed is the Chinese hamster ovary (CHO) cell line deficient in DHFR activity, prepared and propagated as described by Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77:4216 [1980]. The transformed cells are then exposed to increased levels of Mtx. This leads to the synthesis of multiple copies of the DHFR gene, and, concomitantly, multiple copies of other DNA comprising the expression vectors, such as the DNA encoding mpl ligand. This amplification technique can be used with any otherwise suitable host, e.g., ATCC No. CCL61 CHO-K1, notwithstanding the presence of endogenous DHFR if, for example, a mutant DHFR gene that is highly resistant to Mtx is employed (EP 117,060). Alternatively, host cells [particularly wild-type hosts that contain endogenous DHFR] transformed or co-transformed with DNA sequences encoding mpl ligand, wild-type DHFR protein, and another selectable marker such as aminoglycoside 3' phosphotransferase (APH) can be selected by cell growth in medium containing a selection agent for the selectable marker such as an aminoglycosidic antibiotic, e.g., kanamycin, neomycin, or G418. See U.S. Patent No. 4,965,199.

A suitable selection gene for use in yeast is the *trp*1 gene present in the yeast plasmid YRp7 (Stinchcomb *et al.*, *Nature*, **282**:39 [1979]; Kingsman *et al.*, *Gene*, **7**:141 [1979]; or Tschemper *et al.*, *Gene*, **10**:157 [1980]). The *trp*1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 (Jones, *Genetics*, **85**:12 [1977]). The presence of the *trp*1 lesion in the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan. Similarly, *Leu2*-deficient yeast strains (ATCC No. 20,622 or 38,626) are complemented by known plasmids bearing the *Leu2* gene.

#### (iv) Promoter Component

Expression and cloning vectors usually contain a promoter that is recognized by the host organism and is operably linked to the *mpl* ligand nucleic acid. Promoters are untranslated sequences located upstream (5') to the start codon of a structural gene (generally within about 100 to 1000 bp) that control the transcription and translation of particular nucleic acid sequence, such as the *mpl* ligand nucleic acid sequence, to which they are operably linked. Such promoters typically fall into two classes, inducible and constitutive. Inducible promoters are promoters that initiate increased levels of transcription from DNA under their control in response to some change in culture conditions, *e.g.*, the presence or absence of a nutrient or a change in

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temperature. At this time a large number of promoters recognized by a variety of potential host cells are well known. These promoters are operably linked to *mpl* ligand encoding DNA by removing the promoter from the source DNA by restriction enzyme digestion and inserting the isolated promoter sequence into the vector. Both the native *mpl* ligand promoter sequence and many heterologous promoters may be used to direct amplification and/or expression of the *mpl* ligand DNA. However, heterologous promoters are preferred, as they generally permit greater transcription and higher yields of expressed *mpl* ligand as compared to the native *mpl* ligand promoter.

Promoters suitable for use with prokaryotic hosts include the ß-lactamase and lactose promoter systems (Chang et al., Nature, 275:615 [1978]; and Goeddel et al., Nature, 281:544 [1979]), alkaline phosphatase, a tryptophan (trp) promoter system (Goeddel, Nucleic Acids Res., 8:4057 [1980] and EP 36,776) and hybrid promoters such as the tac promoter (deBoer et al., Proc. Natl. Acad. Sci. USA, 80:21-25 [1983]). However, other known bacterial promoters are suitable. Their nucleotide sequences have been published, thereby enabling a skilled worker operably to ligate them to DNA encoding mpl ligand (Siebenlist et al., Cell, 20:269 [1980]) using linkers or adaptors to supply any required restriction sites. Promoters for use in bacterial systems also will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding mpl ligand polypeptide.

Promoter sequences are known for eukaryotes. Virtually all eukaryotic genes have an AT-rich region located approximately 25 to 30 bases upstream from the site where transcription is initiated. Another sequence found 70 to 80 bases upstream from the start of transcription of many genes is a CXCAAT region where X may be any nucleotide. At the 3' end of most eukaryotic genes is an AATAAA sequence that may be the signal for addition of the poly A tail to the 3' end of the coding sequence. All of these sequences are suitably inserted into eukaryotic expression vectors.

Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem., 255:2073 [1980]) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg., 7:149 [1968]; and Holland, Biochemistry, 17:4900 [1978]), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate

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dehydrogenase, and enzymes responsible for maltose and galactose utilization. Sultable vectors and promoters for use in yeast expression are further described in Hitzeman et al., EP 73,657A. Yeast enhancers also are advantageously used with yeast promoters.

MpI ligand transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and most preferably Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, from heat-shock promoters, and from the promoter normally associated with the mpI ligand sequence, provided such promoters are compatible with the host cell systems.

The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment that also contains the SV40 viral origin of replication. Fiers et al., Nature, 273:113 [1978]; Mulligan and Berg, Science, 209:1422-1427 [1980]; Pavlakis et al., Proc. Natl. Acad. Sci. USA, 78:7398-7402 [1981]. The immediate early promoter of the human cytomegalovirus is conveniently obtained as a HindIII E restriction fragment. Greenaway et al., Gene, 18:355-360 [1982]. A system for expressing DNA in mammalian hosts using the bovine papilloma virus as a vector is disclosed in U.S. Patent No. 4,419,446. A modification of this system is described in U.S. Patent No. 4,601,978. See also Gray et al., Nature, 295:503-508 [1982] on expressing cDNA encoding immune interferon in monkey cells; Reyes et al., Nature, 297:598-601 [1982] on expression of human B-interferon cDNA in mouse cells under the control of a thymidine kinase promoter from herpes simplex virus; Canaani and Berg, Proc. Natl. Acad. Sci. USA, 79:5166-5170 [1982] on expression of the human interferon B1 gene in cultured mouse and rabbit cells; and Gorman et al., Proc. Natl. Acad. Sci. USA, 79:6777-6781 [1982] on expression of bacterial CAT sequences in CV-1 monkey kidney cells, chicken embryo fibroblasts, Chinese hamster ovary cells, HeLa cells, and mouse NiH-3T3 cells using the Rous sarcoma virus long terminal repeat as a promoter.

#### (v) Enhancer Element Component

Transcription of a DNA encoding the *mpl* ligand of this invention by higher eukaryotes is often increased by inserting an enhancer sequence into the vector. Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp, that act on a promoter to increase its transcription. Enhancers are relatively orientation and position independent, having been found 5' (Laimins *et al.*, *Proc. Natl. Acad. Sci. USA*, **78**:993 [1981]) and 3' (Lusky *et al.*, *Mol. Cell Bio.*, 3:1108 [1983]) to the

transcription unit, within an intron (Banerji et al., Cell, 33:729 [1983]), as well as within the coding sequence itself (Osborne et al., Mol. Cell Bio., 4:1293 [1984]). Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, a-fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. See also Yaniv, Nature, 297:17-18 [1982] on enhancing elements for activation of eukaryotic promoters. The enhancer may be spliced into the vector at a position 5' or 3' to the mpl ligand encoding sequence, but is preferably located at a site 5' from the promoter.

## (vi) Transcription Termination Component

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Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellular organisms) will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3' untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding *mpl* ligand.

## (vii) Construction and Analysis of Vectors

Construction of suitable vectors containing one or more of the above listed components employs standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to generate the plasmids required.

For analysis to confirm correct sequences in plasmids constructed, the ligation mixtures are used to transform *E. coli* K12 strain 294 (ATCC No. 31,446) and successful transformants selected by ampicillin or tetracycline resistance where appropriate. Plasmids from the transformants are prepared, analyzed by restriction endonuclease digestion, and/or sequenced by the method of Messing *et al.*, *Nucleic Acids Res.*, 9:309 [1981] or by the method of Maxam *et al.*, *Methods in Enzymology*, 65:499 [1980].

## (viii) Transient Expression Vectors

Particularly useful in the practice of this invention are expression vectors that provide for the transient expression in mammalian cells of DNA encoding the *mpl* ligand polypeptide. In general, transient expression involves the use of an expression vector that is able to replicate efficiently in a host cell, such that the host cell accumulates many copies of the expression vector and, in turn, synthesizes high levels of a desired polypeptide encoded by the expression vector. Sambrook *et al.*, *supra*, pp.

16.17 - 16.22. Transient expression systems, comprising a suitable expression vector and a host cell, allow for the convenient positive identification of polypeptides encoded by cloned DNAs, as well as for the rapid screening of such polypeptides for desired biological or physiological properties. Thus, transient expression systems are particularly useful in the invention for purposes of identifying analogues and variants of *mpl* ligand polypeptide that have *mpl* ligand polypeptide biological activity.

#### (ix) Suitable Exemplary Vertebrate Cell Vectors

Other methods, vectors, and host cells suitable for adaptation to the synthesis of *mpl* ligand in recombinant vertebrate cell culture are described in Gething *et al.*, *Nature*, 293:620-625 [1981]; Mantei *et al.*, *Nature*, 281:40-46 [1979]; Levinson *et al.*; EP 117,060; and EP 117,058. A particularly useful plasmid for mammalian cell culture expression of *mpl* ligand is pRK5 (EP 307,247 U. S. patent no. 5,258,287) or pSVI6B (PCT Publication No. WO 91/08291).

#### D. Selection and Transformation of Host Cells

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Sultable host cells for cloning or expressing the vectors herein are the prokaryote, yeast, or higher eukaryotic cells described above. Suitable prokaryotes include eubacteria, such as Gram-negative or Gram-positive organisms, for example, *E. coli*, *Bacilli* such as *B. subtilis*, *Pseudomonas* species such as *P. aeruginosa*, *Salmonella typhimurium*, or *Serratia marcescans*. One preferred *E. coli* cloning host is *E. coli* 294 (ATCC No. 31,446), although other strains such as *E. coli* B, *E. coli* X1776 (ATCC No. 31,537), and *E. coli* W3110 (ATCC No. 27,325) are suitable. These examples are illustrative rather than limiting. Preferably the host cell should secrete minimal amounts of proteolytic enzymes. Alternatively, *in vitro* methods of cloning, *e.g.*, PCR or other nucleic acid polymerase reactions, are suitable.

In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable hosts for *mpl* ligand encoding vectors. Saccharomyces cerevisiae, or common baker's yeast, is the most commonly used among lower eukaryotic host microorganisms. However, a number of other genera, species, and strains are commonly available and useful herein, such as Schizosaccharomyces pombe (Beach and Nurse, Nature, 290:140 [1981]; EP 139,383 published 2 May 1985), Kluyveromyces hosts (U.S. Patent No. 4,943,529) such as, e.g., K. lactis (Louvencourt et al., J. Bacteriol., 737 [1983]), K. fragilis, K. bulgaricus, K. thermotolerans, and K. marxianus, yarrowia [EP 402,226], Pichia pastoris (EP 183,070; Sreekrishna et al., J. Basic Microbiol., 28:265-278 [1988]), Candida, Trichoderma reesia (EP 244,234), Neurospora crassa (Case et al., Proc. Natl. Acad. Sci. USA, 76:5259-5263 [1979]), and filamentous fungi such as, e.g, Neurospora, Penicillium, Tolypocladium (WO 91/00357 published 10 January 1991), and Aspergillus hosts such as A. nidulans (Ballance et al., Biochem. Biophys. Res.

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Commun., 112:284-289 [1983]; Tilburn et al., Gene, 26:205-221 [1983]; Yelton et al., Proc. Natl. Acad. Sci. USA, 81:1470-1474 [1984]) and A. niger (Kelly and Hynes, EMBO J., 4:475-479 [1985]).

Suitable host cells for the expression of glycosylated *mpl* ligand are derived from multicellular organisms. Such host cells are capable of complex processing and glycosylation activities. In principle, any higher eukaryotic cell culture is workable, whether from vertebrate or invertebrate culture. Examples of invertebrate cells include plant and insect cells. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts such as *Spodoptera frugiperda* (caterpillar), *Aedes aegypti* (mosquito), *Aedes albopictus* (mosquito), *Drosophila melanogaster* (fruitfly), and *Bombyx mori* have been identified. See, *e.g.*, Luckow *et al.*, *Bio/Technology*, 6:47-55 [1988]; Miller *et al.*, *Genetic Engineering*, Setlow *et al.*, eds., Vol. 8 (Plenum Publishing, 1986), pp. 277-279; and Maeda *et al.*, *Nature*, 315:592-594 [1985]. A variety of viral strains for transfection are publicly available, *e.g.*, the L-1 variant of *Autographa californica* NPV and the Bm-5 strain of *Bombyx mori* NPV, and such viruses may be used as the virus herein according to the present invention, particularly for transfection of *Spodoptera frugiperda* cells.

Plant cell cultures of cotton, corn, potato, soybean, petunia, tomato, and tobacco can be utilized as hosts. Typically, plant cells are transfected by incubation with certain strains of the bacterium Agrobacterium tumefaciens, which has been previously manipulated to contain the mpl ligand DNA. During incubation of the plant cell culture with A. tumefaciens, the DNA encoding the mpl ligand is transferred to the plant cell host such that it is transfected, and will, under appropriate conditions, express the mpl ligand DNA. In addition, regulatory and signal sequences compatible with plant cells are available, such as the nopaline synthase promoter and polyadenylation signal sequences. Depicker et al., J. Mol. Appl. Gen., 1:561 [1982]. In addition, DNA segments isolated from the upstream region of the T-DNA 780 gene are capable of activating or increasing transcription levels of plant-expressible genes in recombinant DNA-containing plant tissue. EP 321,196 published 21 June 1989.

However, interest has been greatest in vertebrate cells, and propagation of vertebrate cells in culture (tissue culture) has become a routine procedure in recent years (Tissue Culture, Academic Press, Kruse and Patterson, editors [1973]). Examples of useful mammalian host cell lines are monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham et al., J. Gen Virol., 36:59 [1977]); baby hamster kidney cells (BHK, ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77:4216 [1980]); mouse sertoli cells (TM4, Mather, Biol. Reprod., 23:243-251 [1980]); monkey

kidney cells (CV1 ATCC CCL 70); African green monkey kidney cells (VERO-76, ATCC CRL-1587); human cervical carcinoma cells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL 34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); mouse mammary tumor (MMT 060562, ATCC CCL51); TRI cells (Mather et al., Annals N.Y. Acad. Sci., 383:44-68 [1982]); MRC 5 cells; FS4 cells; and a human hepatoma line (Hep G2).

Host cells are transfected and preferably transformed with the above-described expression or cloning vectors of this invention and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences.

Transfection refers to the taking up of an expression vector by a host cell whether or not any coding sequences are in fact expressed. Numerous methods of transfection are known to the ordinarily skilled artisan, for example, CaPO<sub>4</sub> and electroporation. Successful transfection is generally recognized when any indication of the operation of this vector occurs within the host cell.

Transformation means introducing DNA into an organism so that the DNA is replicable, either as an extrachromosomal element or by chromosomal integrant. Depending on the host cell used, transformation is done using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in section 1.82 of Sambrook et al., supra, is generally used for prokaryotes or other cells that contain substantial cell-wall barriers. Infection with Agrobacterium tumefaciens is used for transformation of certain plant cells, as described by Shaw et al., Gene, 23:315 [1983] and WO 89/05859 published 29 June 1989. In addition, plants may be transfected using ultrasound treatment as described in WO 91/00358 published 10 January 1991. For mammalian cells without such cell walls, the calcium phosphate precipitation method of Graham and van der Eb. Virology, 52:456-457 [1978] is preferred. General aspects of mammalian cell host system transformations have been described by Axel in U.S. Patent No. 4,399,216 issued 16 August 1983. Transformations into yeast are typically carried out according to the method of Van Solingen et al., J. Bact., 130:946 [1977] and Hsiao et al., Proc. Natl. Acad. Sci. (USA), 76:3829 [1979]. However, other methods for introducing DNA into cells such as by nuclear injection, electroporation, or protoplast fusion may also be used.

#### E. Culturing the Host Cells

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Prokaryotic cells used to produce the *mpl* ligand polypeptide of this invention are cultured in suitable media as described generally in Sambrook *et al.*, *supra*.

The mammalian host cells used to produce the *mpl* ligand of this invention may be cultured in a variety of media. Commercially available media such as Ham's F10

(Sigma), Minimal Essential Medium ([MEM], Sigma), RPMI-1640 (Sigma), and Dulbecco's Modified Eagle's Medium ([DMEM], Sigma) are suitable for culturing the host cells. In addition, any of the media described in Ham and Wallace, Meth. Enz., 58:44 [1979], Barnes and Sato, Anal. Biochem., 102:255 [1980], U.S. Patent No. 4,767,704; 4,657,866; 4,927,762; or 4,560,655; WO 90/03430; WO 87/00195; U.S. Patent Re. 30,985; or copending U.S.S.N. 07/592,107 or 07/592,141, both filed on 3 October 1990, the disclosures of all of which are incorporated herein by reference, may be used as culture media for the host cells. Any of these media may be supplemented as necessary with hormones and/or other growth factors (such as insulin, transferrin, or epidermal growth factor), salts (such as sodium chloride, calcium, magnesium, and phosphate), buffers (such as HEPES), nucleosides (such as adenosine and thymidine), antibiotics (such as Gentamycin $^{\text{TM}}$ drug), trace elements (defined as inorganic compounds usually present at final concentrations in the micromolar range), and glucose or an equivalent energy source. Any other necessary supplements may also be included at appropriate concentrations that would be known to those skilled in the art. The culture conditions, such as temperature, pH, and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

The host cells referred to in this disclosure encompass cells in in vitro culture as well as cells that are within a host animal.

## F. Detecting Gene Amplification/Expression

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Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, northern blotting to quantitate the transcription of mRNA (Thomas, *Proc. Natl. Acad. Sci. USA*, 77:5201-5205 [1980]), dot blotting (DNA analysis), or *in situ* hybridization, using an appropriately labeled probe, based on the sequences provided herein. Various labels may be employed, most commonly radioisotopes, particularly <sup>32</sup>P. However, other techniques may also be employed, such as using biotin-modified nucleotides for introduction into a polynucleotide. The biotin then serves as the site for binding to avidin or antibodies, which may be labeled with a wide variety of labels, such as radionuclides, fluorescers, enzymes, or the like. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining of tissue sections and assay of cell culture or

body fluids, to quantitate directly the expression of gene product. With immunohistochemical staining techniques, a cell sample is prepared, typically by dehydration and fixation, followed by reaction with labeled antibodies specific for the gene product coupled, where the labels are usually visually detectable, such as enzymatic labels, fluorescent labels, luminescent labels, and the like. A particularly sensitive staining technique suitable for use in the present invention is described by Hsu et al., Am. J. Clin. Path., 75:734-738 [1980].

Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native *mpl* ligand polypeptide or against a synthetic peptide based on the DNA sequences provided herein as described further below.

#### G. Purification of mpl ligand Polypeptide

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*Mpl* ligand preferably is recovered from the culture medium as a secreted polypeptide, although it also may be recovered from host cell lysates when directly expressed without a secretory signal.

When mpl ligand is expressed in a recombinant cell other than one of human origin, the mpl ligand is completely free of proteins or polypeptides of human origin. However, it is still usually necessary to purify mpl ligand from other recombinant cell proteins or polypeptides to obtain preparations that are substantially homogeneous as to the mpl ligand per se. As a first step, the culture medium or lysate is centrifuged to remove particulate cell debris. The membrane and soluble protein fractions are then separated. Alternatively, a commercially available protein concentration filter (e.g., Amicon or Millipore Pellicon ultrafiltration units) may be used. The mpl ligand may then be purified from the soluble protein fraction and from the membrane fraction of the culture lysate, depending on whether the mpl ligand is membrane bound. Mpl ligand thereafter is purified from contaminant soluble proteins and polypeptides by salting out and exchange or chromatographic procedures employing various gel matrices. These matrices include; acrylamide, agarose, dextran, cellulose and others common to protein purification. Exemplary chromatography procedures suitable for protein purification include; immunoaffinity (e.g., anti-hmpl ligand Mab), receptoraffinity (e.g., mpl-lgG or protein A Sepharose), hydrophobic interaction chromatography (HIC) (e.g., ether, butyl, or phenyl Toyopearl), lectin chromatography (e.g., Con A-Sepharose, lentil-lectin-Sepharose), size exclusion (e.g., Sephadex G-75), cation- and anion-exchange columns (e.g., DEAE or carboxymethyl- and sulfopropyl-cellulose), and reverse-phase high performance liquid chromatography (RP-HPLC) (see e.g., Urdal et al., J. Chromatog., 296:171 [1984] where two sequential RP-HPLC steps are used to purify recombinant human

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IL-2). Other purification steps optionally include; ethanol precipitation; ammonium sulfate precipitation; chromatofocusing; preparative SDS-PAGE, and the like.

Mpl ligand variants in which residues have been deleted, inserted, or substituted are recovered in the same fashion as native mpl ligand, taking account of any substantial changes in properties occasioned by the variation. For example, preparation of a mpl ligand fusion with another protein or polypeptide, e.g., a bacterial or viral antigen, facilitates purification; an immunoaffinity column containing antibody to the antigen can be used to adsorb the fusion polypeptide. Immunoaffinity columns such as a rabbit polyclonal anti-mpl ligand column can be employed to absorb the mpl ligand variant by binding it to at least one remaining Alternatively, the mpl ligand may be purified by affinity immune epitope. chromatography using a purified mpl-lgG coupled to a (preferably) immobilized resin such as Affi-Gel 10 (Bio-Rad, Richmond, CA) or the like, by means well known in the art. A protease inhibitor such as phenyl methyl sulfonyl fluoride (PMSF) also may be useful to inhibit proteolytic degradation during purification, and antibiotics may be included to prevent the growth of adventitious contaminants. One skilled in the art will appreciate that purification methods suitable for native mpl ligand may require modification to account for changes in the character of mpl ligand or its variants upon expression in recombinant cell culture.

## H. Covalent Modifications of mpl ligand Polypeptide

Covalent modifications of *mpl* ligand polypeptides are included within the scope of this invention. Both native *mpl* ligand and amino acid sequence variants of the *mpl* ligand may be covalently modified. One type of covalent modification included within the scope of this invention is a *mpl* ligand fragment. Variant *mpl* ligand fragments having up to about 40 amino acid residues may be conveniently prepared by chemical synthesis or by enzymatic or chemical cleavage of the full-length or variant *mpl* ligand polypeptide. Other types of covalent modifications of the *mpl* ligand or fragments thereof are introduced into the molecule by reacting targeted amino acid residues of the *mpl* ligand or fragments thereof with an organic derivatizing agent that is capable of reacting with selected side chains or the N- or C-terminal residues.

Cysteinyl residues most commonly are reacted with  $\alpha$ -haloacetates (and corresponding amines), such as chloroacetic acid or chloroacetamide, to give carboxymethyl or carboxyamidomethyl derivatives. Cysteinyl residues also are derivatized by reaction with bromotrifluoroacetone,  $\alpha$ -bromo- $\beta$ -(5-imidozoyl)propionic acid, chloroacetyl phosphate, N-alkylmaleimides, 3-nitro-2-pyridyl disulfide, methyl 2-pyridyl disulfide, p-chloromercuribenzoate, 2-chloromercuri-4-nitrophenol, or chloro-7-nitrobenzo-2-oxa-1,3-diazole.

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Histidyl residues are derivatized by reaction with diethylpyrocarbonate at pH 5.5-7.0 because this agent is relatively specific for the histidyl side chain. Parabromophenacyl bromide also is useful; the reaction is preferably performed in 0.1M sodium cacodylate at pH 6.0.

Lysinyl and amino terminal residues are reacted with succinic or other carboxylic acid anhydrides. Derivatization with these agents has the effect of reversing the charge of the lysinyl residues. Other suitable reagents for derivatizing -amino-containing residues include imidoesters such as methyl picolinimidate; pyridoxal phosphate; pyridoxal; chloroborohydride; trinitrobenzenesulfonic acid; Omethylisourea; 2,4-pentanedione; and transaminase-catalyzed reaction with glyoxylate.

Arginyl residues are modified by reaction with one or several conventional reagents, among them phenylglyoxal, 2,3-butanedione, 1,2-cyclohexanedione, and ninhydrin. Derivatization of arginine residues requires that the reaction be performed in alkaline conditions because of the high  $pK_a$  of the guanidine functional group. Furthermore, these reagents may react with the groups of lysine as well as the arginine epsilon-amino group.

The specific modification of tyrosyl residues may be made, with particular interest in introducing spectral labels into tyrosyl residues by reaction with aromatic diazonium compounds or tetranitromethane. Most commonly, N-acetylimidizole and tetranitromethane are used to form O-acetyl tyrosyl species and 3-nitro derivatives, respectively. Tyrosyl residues are iodinated using <sup>125</sup>I or <sup>131</sup>I to prepare labeled proteins for use in radioimmunoassay, the chloramine T method described above being suitable.

Carboxyl side groups (aspartyl or glutamyl) are selectively modified by reaction with carbodiimides (R-N=C=N-R'), where R and R' are different alkyl groups, such as 1-cyclohexyl-3-(2-morpholinyl-4-ethyl)carbodiimide or 1-ethyl-3-(4-azonia-4,4-dimethylpentyl)carbodiimide. Furthermore, aspartyl and glutamyl residues are converted to asparaginyl and glutaminyl residues by reaction with ammonium ions.

Derivatization with bifunctional agents is useful for crosslinking *mpl* ligand to a water-insoluble support matrix or surface for use in the method for purifying anti-*mpl* ligand antibodies, and *vice versa*. Commonly used crosslinking agents include, *e.g.*,
1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide
esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters,
including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), and
bifunctional maleimides such as bis-N-maleimido-1,8-octane. Derivatizing agents
such as methyl-3-[(p-azidophenyl)dithio]propioimidate yield photoactivatable

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intermediates that are capable of forming crosslinks in the presence of light. Alternatively, reactive water-insoluble matrices such as cyanogen bromide-activated carbohydrates and the reactive substrates described in U.S. Patent Nos. 3,969,287; 3,691,016; 4,195,128; 4,247,642; 4,229,537; and 4,330,440 are employed for protein immobilization.

Glutaminyl and asparaginyl residues are frequently deamidated to the corresponding glutamyl and aspartyl residues, respectively. These residues are deamidated under neutral or basic conditions. The deamidated form of these residues falls within the scope of this invention.

Other modifications include hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the  $\alpha$ -amino groups of lysine, arginine, and histidine side chains (T.E. Creighton, *Proteins: Structure and Molecular Properties*, W.H. Freeman & Co., San Francisco, pp. 79-86 [1983]), acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the *mpl* ligand polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. By altering is meant deleting one or more carbohydrate moieties found in native *mpl* ligand, and/or adding one or more glycosylation sites that are not present in the native *mpl* ligand.

Glycosylation of polypeptides is typically either N-linked or O-linked. N-linked refers to the attachment of the carbohydrate moiety to the side chain of an asparagine residue. The tripeptide sequences asparagine-X-serine and asparagine-X-threonine, where X is any amino acid except proline, are the recognition sequences for enzymatic attachment of the carbohydrate moiety to the asparagine side chain. Thus, the presence of either of these tripeptide sequences in a polypeptide creates a potential glycosylation site. O-linked glycosylation refers to the attachment of one of the sugars N-acetylgalactosamine, galactose, or xylose to a hydroxyamino acid, most commonly serine or threonine, although 5-hydroxyproline or 5-hydroxylysine may also be used.

Addition of glycosylation sites to the *mpl* ligand polypeptide is conveniently accomplished by altering the amino acid sequence such that it contains one or more of the above-described tripeptide sequences (for N-linked glycosylation sites). The alteration may also be made by the addition of, or substitution by, one or more serine or threonine residues to the native *mpl* ligand sequence (for O-linked glycosylation sites). For ease, the *mpl* ligand amino acid sequence is preferably altered through changes at the DNA level, particularly by mutating the DNA encoding the *mpl* ligand polypeptide at preselected bases such that codons are generated that will translate into

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the desired amino acids. The DNA mutation(s) may be made using methods described above under the heading of "Amino Acid Sequence Variants of mpl Ligand."

Another means of increasing the number of carbohydrate moieties on the *mpl* ligand is by chemical or enzymatic coupling of glycosides to the polypeptide. These procedures are advantageous in that they do not require production of the polypeptide in a host cell that has glycosylation capabilities for N- or O-linked glycosylation. Depending on the coupling mode used, the sugar(s) may be attached to (a) arginine and histidine, (b) free carboxyl groups, (c) free sulfhydryl groups such as those of cysteine, (d) free hydroxyl groups such as those of serine, threonine, or hydroxyproline, (e) aromatic residues such as those of phenylalanine, tyrosine, or tryptophan, or (f) the amide group of glutamine. These methods are described in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, *CRC Crit. Rev. Biochem.*, pp. 259-306 [1981].

Removal of carbohydrate moieties present on the *mpl* ligand polypeptide may be accomplished chemically or enzymatically. Chemical deglycosylation requires exposure of the polypeptide to the compound trifluoromethanesulfonic acid, or an equivalent compound. This treatment results in the cleavage of most or all sugars except the linking sugar (N-acetylglucosamine or N-acetylgalactosamine), while leaving the polypeptide intact. Chemical deglycosylation is described by Hakimuddin, *et al.*, *Arch. Biochem. Biophys.*, **259**:52 [1987] and by Edge *et al.*, *Anal. Biochem.*, **118**:131 [1981]. Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo- and exo-glycosidases as described by Thotakura *et al.*, *Meth. Enzymol.*, **138**:350 [1987].

Glycosylation at potential glycosylation sites may be prevented by the use of the compound tunicamycin as described by Duskin *et al.*, *J. Biol. Chem.*, **257**:3105 [1982]. Tunicamycin blocks the formation of protein-N-glycoside linkages.

Another type of covalent modification of *mpl* ligand comprises linking the *mpl* ligand polypeptide to one of a variety of nonproteinaceous polymers, *e.g.*, polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337. *Mpl* ligand polypeptides covalently linked to the forgoing polymers are referred to herein as pegylated *mpl* ligand polypeptides

It will be appreciated that some screening of the recovered *mpl* ligand variant will be needed to select the optimal variant for binding to a *mpl* and having the immunological and/or biological activity defined above. One can screen for stability in recombinant cell culture or in plasma (e.g., against proteolytic cleavage), high affinity to a *mpl* member, oxidative stability, ability to be secreted in elevated yields, and the like. For example, a change in the immunological character of the *mpl* ligand

polypeptide, such as affinity for a given antibody, is measured by a competitive-type immunoassay. Other potential modifications of protein or polypeptide properties such as redox or thermal stability, hydrophobicity, or susceptibility to proteolytic degradation are assayed by methods well known in the art.

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# 17. General Methods for Preparation of Antibodies to mpl Ligand Antibody Preparation

## (i) Polyclonal antibodies

Polyclonal antibodies to *mpl* ligand polypeptides or fragments are generally raised in animals by multiple subcutaneous (sc) or intraperitoneal (ip) injections of the *mpl* ligand and an adjuvant. It may be useful to conjugate the *mpl* ligand or a fragment containing the target amino acid sequence to a protein that is immunogenic in the species to be immunized, *e.g.*, keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, or soybean trypsin inhibitor using a bifunctional or derivatizing agent, for example maleimidobenzoyl sulfosuccinimide ester (conjugation through cysteine residues), N-hydroxysuccinimide (through lysine residues), glytaraldehyde, succinic anhydride, SOCl<sub>2</sub>, or R<sup>1</sup>N=C=NR, where R and R<sup>1</sup> are different alkyl groups.

Animals are immunized against the *mpl* ligand polypeptide or fragment, immunogenic conjugates or derivatives by combining 1 mg of 1 µg of the peptide or conjugate (for rabbits or mice, respectively) with 3 volumes of Freund's complete adjuvant and injecting the solution intradermally at multiple sites. One month later the animals are boosted with 1/5 to 1/10 the original amount of peptide or conjugate in Freund's complete adjuvant by subcutaneous injection at multiple sites. Seven to 14 days later the animals are bled and the serum is assayed for *mpl* ligand antibody titer. Animals are boosted until the titer plateaus. Preferably, the animal boosted with the conjugate of the same *mpl* ligand, but conjugated to a different protein and/or through a different cross-linking reagent. Conjugates also can be made in recombinant cell culture as protein fusions. Also, aggregating agents such as alum are used to enhance the immune response.

## (ii) Monoclonal antibodies

Monoclonal antibodies are obtained from a population of substantially homogeneous antibodies, *i.e.*, the individual antibodies comprising the population are identical except for possible naturally occurring mutations that may be present in minor amounts. Thus, the modifier "monoclonal" indicates the character of the antibody as not being a mixture of discrete antibodies.

For example, the mpl ligand monoclonal antibodies of the invention may be made using the hybridoma method first described by Kohler & Milstein, Nature,

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256:495 [1975], or may be made by recombinant DNA methods (U.S. Patent No. 4,816,567 [Cabilly *et al.*]).

In the hybridoma method, a mouse or other appropriate host animal, such as hamster is immunized as hereinabove described to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the protein used for immunization. Alternatively, lymphocytes may be immunized *in vitro*. Lymphocytes then are fused with myeloma cells using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, *Monoclonal Antibodies: Principles and Practice*, pp.59-103 [Academic Press, 1986]).

The hybridoma cells thus prepared are seeded and grown in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells. For example, if the parental myeloma cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substances prevent the growth of HGPRT-deficient cells.

Preferred myeloma cells are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. Among these, preferred myeloma cell lines are murine myeloma lines, such as those derived from MOPC-21 and MPC-11 mouse tumors available from the Salk Institute Cell Distribution Center, San Diego, California USA, and SP-2 cells available from the American Type Culture Collection, Rockville, Maryland USA. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 [1984]; Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp.51-63, Marcel Dekker, Inc., New York, 1987).

Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against *mpl* ligand. Preferably, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an *in vitro* binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA).

The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson & Pollard, *Anal. Biochem.*, **107**:220 [1980].

After hybridoma cells are identified that produce antibodies of the desired specificity, affinity, and/or activity, the clones may be subcloned by limiting dilution procedures and grown by standard methods (Goding, *supra*). Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium or RPMI-

1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in an animal.

The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

DNA encoding the monoclonal antibodies of the invention is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences, (Cabilly et al., supra; Morrison, et al., Proc. Nat. Acad. Sci., 81:6851 [1984]), or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide.

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Typically such non-immunoglobulin polypeptides are substituted for the constant domains of an antibody of the invention, or they are substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody comprising one antigen-combining site having specificity for a *mpl* ligand and another antigen-combining site having specificity for a different antigen.

Chimeric or hybrid antibodies also may be prepared *in vitro* using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins may be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate.

For diagnostic applications, the antibodies of the invention typically will be labeled with a detectable moiety. The detectable moiety can be any one which is capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety may be a radioisotope, such as <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, or <sup>125</sup>I, a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin; radioactive isotopic labels, such as, *e.g.*, <sup>125</sup>I, <sup>32</sup>P, <sup>14</sup>C, or <sup>3</sup>H, or an enzyme, such as alkaline phosphatase, beta-galactosidase or horseradish peroxidase.

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Any method known in the art for separately conjugating the antibody to the detectable molety may be employed, including those methods described by Hunter, et al., Nature, 144:945 [1962]; David, et al., Biochemistry, 13:1014 [1974]; Pain, et al., J. Immunol. Meth., 40:219 [1981]; and Nygren, J. Histochem. and Cytochem., 30:407 [1982].

The antibodies of the present invention may be employed in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays. Zola, *Monoclonal Antibodies: A Manual of Techniques*, pp.147-158 (CRC Press, Inc., 1987).

Competitive binding assays rely on the ability of a labeled standard (which may be a *mpl* ligand or an immunologically reactive portion thereof) to compete with the test sample analyte (*mpl* ligand) for binding with a limited amount of antibody. The amount of *mpl* ligand in the test sample is inversely proportional to the amount of standard that becomes bound to the antibodies. To facilitate determining the amount of standard that becomes bound, the antibodies generally are insolubilized before or after the competition, so that the standard and analyte that are bound to the antibodies may conveniently be separated from the standard and analyte which remain unbound.

Sandwich assays involve the use of two antibodies, each capable of binding to a different immunogenic portion, or epitope, of the protein (*mpl* ligand) to be detected. In a sandwich assay, the test sample analyte is bound by a first antibody which is immobilized on a solid support, and thereafter a second antibody binds to the analyte, thus forming an insoluble three part complex. David & Greene, U.S. Patent No. 4,376,110. The second antibody may itself be labeled with a detectable moiety (direct sandwich assays) or may be measured using an anti-immunoglobulin antibody that is labeled with a detectable moiety (indirect sandwich assay). For example, one type of sandwich assay is an ELISA assay, in which case the detectable moiety is an enzyme (e.g., horseradish peroxidase).

#### (iii) Humanized and human antibodies

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers (Jones et al., Nature, 321:522-525 [1986]; Riechmann et al., Nature, 332:323-327 [1988]; Verhoeyen et al., Science, 239:1534-1536 [1988]), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (Cabilly et al., supra), wherein substantially less than an intact human

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variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies is very important in order to reduce antigenicity. According to the so called "best-fit" method, the sequence of the variable domain of a rodent antibody is screened against the entire library of known human variable domain sequences. The human sequence which is closest to that of the rodent is then accepted as the human framework (FR) for the humanized antibody (Sims et al., J. Immunol., 151:2296 [1993]; Chothia and Lesk, J. Mol. Biol., 196:901 [1987]). Another method uses a particular framework derived from the concensus sequence of all human antibodies of a particular subgroup of light or heavy chains. The same framework may be used for several different humanized antibodies (Carter et al., Proc. Natl. Acad. Sci. USA, 89:4285 [1992]; Presta et al., J. Immnol., 151:2623 [1993]).

It is further important that antibodies be humanized with retention of high. affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three dimensional models of the parental and humanized sequences. Three dimensional immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable threedimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the consensus and import sequence so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. In general, the CDR residues are directly and most substantially involved in influencing antigen binding. For further details see U.S. application Serial No. 07/934,373 filed 21 August 1992, which is a continuationin-part of application Serial No. 07/715,272 filed 14 June 1991.

Alternatively, it is now possible to produce transgenic animals (e.g., mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production. For example, it has been described that the homozygous deletion of the antibody heavy chain joining region ( $J_H$ ) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line

immunoglobulin gene array in such germ-line mutant mice will result in the production of human antibodies upon antigen challenge. See, e.g., Jakobovits et al., Proc. Natl. Acad. Sci. USA, 90:2551-255 [1993]; Jakobovits et al., Nature, 362:255-258 [1993]; Bruggermann et al., Year in Immuno., 7:33 [1993]. Human antibodies can also be produced in phage display libraries (Hoogenboom and Winter, J. Mol. Biol. 227, 381 [1991]; Marks et al., J. Mol. Biol. 222, 581 [1991]).

#### (iv) Bispecific antibodies

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Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. Methods for making bispecific antibodies are known in the art.

Traditionally, the recombinant production of bispecific antibodies is based on the coexpression of two immunoglobulin heavy chain-light chain pairs, where the two heavy chains have different specificities (Millstein and Cuello, *Nature*, **305**:537-539 [1983]). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of 10 different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule, which is usually done by affinity chromatography steps, is rather cumbersome, and the product yields are low. Similar procedures are disclosed in PCT publication No. WO 93/08829 (published 13 May 1993), and in Traunecker *et al.*, *EMBO*, **10**:3655-3659 [1991].

According to a different and more preferred approach, antibody variable domains with the desired binding specificities (antibody-antigen combining sites) are fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy chain constant domain, comprising at least part of the hinge, CH2 and CH3 regions. It is preferred to have the first heavy chain constant region (CH1) containing the site necessary for light chain binding, present in at least one of the fusions. DNAs encoding the immunoglobulin heavy chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are cotransfected into a suitable host organism. This provides for great flexibility in adjusting the mutual proportions of the three polypeptide fragments in embodiments when unequal ratios of the three polypeptide chains used in the construction provide the optimum yields. It is, however, possible to insert the coding sequences for two or all three polypeptide chains in one expression vector when the expression of at least two polypeptide chains in equal ratios results in high yields or when the ratios are of no particular significance. In a preferred embodiment of this approach, the bispecific antibodies are composed of a hybrid immunoglobulin heavy chain with a first binding specificity in one arm, and a hybrid immunoglobulin heavy chain-light chain pair (providing a second binding specificity) in the other arm. It was found that this

asymmetric structure facilitates the separation of the desired bispecific compound from unwanted immunoglobulin chain combinations, as the presence of an immunoglobulin light chain in only one half of the bispecific molecule provides for a facile way of separation. This approach is disclosed in copending application Serial No. 07/931,811 filed 17 August 1992.

For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 [1986].

## (v) Heteroconjugate antibodies

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Heteroconjugate antibodies are also within the scope of the present invention.

Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (U.S. Patent No. 4,676,980), and for treatment of HIV infection (PCT publication Nos. WO 91/00360 and WO 92/00373; EP 03089). Heteroconjugate antibodies may be made using any convenient cross-linking methods. Suitable cross-linking agents are well known in the art, and are disclosed in U.S. Patent No. 4,676,980, along with a number of cross-linking techniques.

## IV. Therapeutic Use of the Megakaryocytopoletic Protein mpl Ligand

The biologically active *mpl* ligand having hematopoietic effector function and referred to here as a megakaryocytopoletic or thrombocytopoletic protein (TPO) may be used in a sterile pharmaceutical preparation or formulation to stimulate megakaryocytopoietic or thrombopoletic activity in patients suffering from thrombocytopenia due to impaired production, sequestration, or increased destruction of platelets. Thrombocytopenia-associated bone marrow hypoplasia (e.g., aplastic anemia following chemotherapy or bone marrow transplant) may be effectively treated with the compounds of this invention as well as disorders such as disseminated intravascular coagulation (DIC), immune thrombocytopenia (including HIV-induced ITP and non HIV-induced ITP), chronic idiopathic thrombocytopenia, congenital thrombocytopenia, myelodysplasia, and thrombotic thrombocytopenia. Additionally, these megakaryocytopoietic proteins may be useful in treating myeloproliferative thrombocytotic diseases as well as thrombocytosis from inflammatory conditions and in Iron deficiency.

Preferred uses of the megakaryocytopoietic or thrombocytopoietic protein (TPO) of this invention are in: myelotoxic chemotherapy for treatment of leukemia or solid tumors, myeloablative chemotherapy for autologous or allogeneic bone marrow transplant, myelodysplasia, idiopathic aplastic anemia, congenital thrombocytopenia, and immune thrombocytopenia.

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Still other disorders usefully treated with the megakaryocytopoietic proteins of this invention include defects or damage to platelets resulting from drugs, poisoning or activation on artificial surfaces. In these cases, the instant compounds may be employed to stimulate "shedding" of new "undamaged" platelets. For a more complete list of useful applications, see the "Background" *supra*, especially section (a)-(f) and references cited therein.

The megakaryocytopoietic proteins of the instant invention may be employed alone or in combination with other cytokines, hematopoletins, interleukins, growth factors, or antibodies in the treatment of the above-identified disorders and conditions. Thus, the instant compounds may be employed in combination with other protein or peptide having thrombopoietic activity including; G-CSF, GM-CSF, LIF, M-CSF, IL-1, IL-3, erythropoietin (EPO), kit ligand, IL-6, and IL-11.

The megakaryocytopoletic proteins of the instant invention are prepared in a mixture with a pharmaceutically acceptable carrier. This therapeutic composition can be administered intravenously or through the nose or lung. The composition may also be administered parenterally or subcutaneously as desired. When administered systematically, the therapeutic composition should be pyrogen-free and in a parenterally acceptable solution having due regard for pH, isotonicity, and stability. These conditions are known to those skilled in the art. Briefly, dosage formulations of the compounds of the present invention are prepared for storage or administration by mixing the compound having the desired degree of purity with physiologically acceptable carriers, excipients, or stabilizers. Such materials are non-toxic to the recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, acetate and other organic acid salts; antioxidants such as ascorbic acid; low molecular weight (less than about ten residues) peptides such as polyarginine, proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidinone; amino acids such as glycine, glutamic acid, aspartic acid, or arginine; monosaccharides, disaccharides, and other carbohydrates including cellulose or its derivatives, glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sobitol; counterions such as sodium and/or nonionic surfactants such as Tween, Pluronics or polyethyleneglycol.

About 0.5 to 500 mg of a compound or mixture of the megakaryocytopoletic protein as the free acid or base form or as a pharmaceutically acceptable salt, is compounded with a physiologically acceptable vehicle, carrier, excipient, binder, preservative, stabilizer, flavor, etc., as called for by accepted pharmaceutical practice. The amount of active ingredient in these compositions is such that a suitable dosage in the range indicated is obtained.

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Sterile compositions for injection can be formulated according to conventional pharmaceutical practice. For example, dissolution or suspension of the active compound in a vehicle such as water or naturally occurring vegetable oil like sesame, peanut, or cottonseed oil or a synthetic fatty vehicle like ethyl oleate or the like may be desired. Buffers, preservatives, antioxidants and the like can be incorporated according to accepted pharmaceutical practice.

Suitable examples of sustained-release preparations include semipermeable matrices of solid hydrophobic polymers containing the polypeptide, which matrices are in the form of shaped articles, e.g., films, or microcapsules. Examples of sustained-release matrices include polyesters, hydrogels [e.g., poly(2-hydroxyethyl-methacrylate) as described by Langer et al., J. Biomed. Mater. Res., 15:167-277 [1981] and Langer, Chem. Tech., 12:98-105 [1982] or poly(vinylalcohol)], polylactides (U.S. Patent No. 3,773,919, EP 58,481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., Biopolymers, 22:547-556 [1983]), non-degradable ethylene-vinyl acetate (Langer et al., supra), degradable lactic acid-glycolic acid copolymers such as the Lupron Depot™ (injectable microspheres composed of lactic acid-glycolic acid copolymer and leuprolide acetate), and poly-D-(-)-3-hydroxybutyric acid (EP 133,988).

While polymers such as ethylene-vinyl acetate and lactic acid-glycolic acid enable release of molecules for over 100 days, certain hydrogels release proteins for shorter time periods. When encapsulated proteins remain in the body for a long time, they may denature or aggregate as a result of exposure to moisture at 37°C, resulting in a loss of biological activity and possible changes in immunogenicity. Rational strategies can be devised for protein stabilization depending on the mechanism involved. For example, if the aggregation mechanism is discovered to be intermolecular S-S bond formation through disulfide interchange, stabilization may be achieved by modifying sulfhydryl residues, lyophilizing from acidic solutions, controlling moisture content, using appropriate additives, and developing specific polymer matrix compositions.

Sustained-release megakaryocytopoietic protein compositions also include liposomally entrapped megakaryocytopoietic protein. Liposomes containing megakaryocytopoietic protein are prepared by methods known per se: DE 3,218,121; Epstein et al., Proc. Natl. Acad. Sci. USA, 82:3688-3692 [1985]; Hwang et al., Proc. Natl. Acad. Sci. USA, 77:4030-4034 [1980]; EP 52,322; EP 36,676; EP 88,046; EP 143,949; EP 142,641; Japanese patent application 83-118008; U.S. Patent Nos. 4,485,045 and 4,544,545; and EP 102,324. Ordinarily the liposomes are of the small (about 200-800 Angstroms) unilamellar type in which the lipid content is

greater than about 30 mol. % cholesterol, the selected proportion being adjusted for the optimal megakaryocytopoietic protein therapy.

The dosage will be determined by the attending physician taking into consideration various factors known to modify the action of drugs including severity and type of disease, body weight, sex, diet, time and route of administration, other medications and other relevant clinical factors. Typically, the daily regimen will range from 0.1-100 µg/kg body weight. Preferably the dosage will range from 0.1-50 µg/kg body weight. More preferably, the initial dosage will range from 1 to 5 µg/kg/day. Optionally, the dosage range will be the same as that of other cytokines, especially G-CSF, GM-CSF, and EPO. Therapeutically effective dosages may be determined by either *in vitro* or *in vivo* methods.

#### **EXAMPLES**

Without further description, it is believed that one of ordinary skill in the art can, using the preceding description and illustrative examples, make and utilize the present invention to the fullest extent. The following working examples therefore specifically point out preferred embodiments of the present invention, and are not to be construed as limiting in any way of the remainder of the disclosure.

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#### **EXAMPLE 1**

#### Partial Purification of the Porcine mpl Ligand

Platelet-poor plasma was collected from normal or aplastic anemic pigs. Pigs were rendered aplastic by irradiation with 900 cGy of total body irradiation using a 4mEV linear accelerator. The irradiated pigs were supported for 6-8 days with intramuscular injections of cefazolin. Subsequently, their total blood volume was removed under general anesthesia, heparinized, and centrifuged at 1800 x g for 30min. to make platelet-poor plasma. The megakaryocyte stimulating activity was found to peak 6 days after irradiation.

Aplastic porcine plasma obtained from irradiated pigs is made 4M with NaCl and stirred for 30 min. at room temperature. The resultant precipitate is removed by centrifugation at 3800 rpm in a Sorvall RC3B and the supernatant is loaded onto a Phenyl-Toyopearl column (220 ml) equilibrated in 10 mM NaPO4 containing 4M NaCl. The column is washed with this buffer until A280 is <0.05 and eluted with dH2O. The eluted protein peak is diluted with dH2O to a conductivity of 15mS and loaded onto a Blue-Sepharose column equilibrated (240 ml) in PBS. Subsequently, the column is washed with 5 column volumes each of PBS and 10mM NaPO4 (pH 7.4) containing 2M urea. Proteins are eluted from the column with 10mM NaPO4 (pH 7.4) containing 2M urea and 1M NaCl. The eluted protein peak is made 0.01% octyl

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glucoside(n-octyl β-D-glucopyranoside) and 1 mM each with EDTA and Pefabloc (Boehinger Mannheim) and loaded directly onto tandemly linked CD4-lgG (Capon, D.J. et al. Nature 337:525-531 [1989]) and mpl-lgG Ultralink (Pierce) columns (see below). The CD4-lgG (2 ml) column is removed after the sample is loaded and the mpl-lgG (4 ml) column is washed with 10 column volumes each of PBS and PBS containing 2 M NaCl and eluted with 0.1M glycine-HCl pH 2.25. Fractions are collected into 1/10th volume 1M Tris-HCl (pH 8.0).

Analysis of eluted fractions from the *mpl*-affinity column by SDS-PAGE (4-20%, Novex gel) run under reducing conditions, revealed the presence of several proteins (Fig. 5). Proteins that silver stain with the strongest intensity resolve with apparent Mr of 66,000, 55,000, 30,000, 28,000 and 14,000. To determine which of these proteins stimulate proliferation of Ba/F3-*mpl* cell cultures these proteins were eluted from the gel as described in Example 2 below.

## Ultralink Affinity Columns

10-20 mg of *mpl*-IgG or CD4-IgG in PBS are coupled to 0.5 grams of Ultralink resin (Pierce) as described by the manufacturer's instructions.

## Construction and Expression of mpl-lgG

A chimeric molecule comprising the entire extracellular domain of human *mpl* (amino acids 1-491) and the Fc region of a human lgG1 molecule was expressed in 293 cells. A cDNA fragment encoding amino acids 1-491 of human *mpl* was obtained by PCR from a human megakaryocytic CMK cell cDNA library and sequenced. A Clal site was inserted at the 5' end and a BstEll site at the 3' end. This fragment was cloned upstream of the lgGl Fc coding region in a Bluescript vector between the Clal and the BstEll sites after partial digestion of the PCR product with BstEll because of two other BstEll sites present in the DNA encoding the extracellular domain of *mpl*. The BstEll site introduced at the 3' end of the *mpl* PCR product was designed to have the Fc region in frame with the *mpl* extracellular domain. The construct was subcloned into pRK5-tkneo vector between the Clal and Xbal sites and transfected into 293 human embryonic kidney cells by the calcium phosphate method. The cells were selected in 0.4 mg/ml G418 and individual clones were isolated. *Mpl*-igG expression from isolated clones was determined using a human Fc specific ELISA. The best expression clone had an expression level of 1-2 mg/ml of *mpl*-lgG.

#### Ba/F3 mpl P Expressing Cells

A cDNA corresponding to the entire coding region of human *mpl* P was cloned into pRK5-tkneo which was subsequently linearized with Notl and transfected into the IL-3 dependent cell line Ba/F3 by electroporation (1 x 10<sup>7</sup> cells, 9605F, 250Volts). Three days later selection was started in the presence of 2 mg/ml of G418. The cells were selected as pools or individual clones were obtained by limiting dilution in 96

well plates. Selected cells were maintained in RPMI containing 15% FBS, 1mg /ml G418, 20mM Glutamine, 10mM HEPES and 100 μg/ml of Pen-Strep. Expression of *mpl* P in selected clones was determined by FACS analysis using a anti-*mpl* P rabbit polyclonal antibody.

Ba/F3 mpl ligand Assay

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The *mpl* ligand assay was conducted as shown is **Fig. 2**. To determine the presence of *mpl* ligand from various sources, the *mpl* P Ba/F3 cells were starved of IL-3 for 24 hours at a cell density of 5 x  $10^5$  cells/ml in a humidified incubator at  $37^{\circ}$ C in 5% CO<sub>2</sub> and air. Following IL-3 starvation the cells were plated out in 96 well culture dishes at a density of 50,000 cells in 200  $\mu$ l of media with or without diluted samples and cultured for 24 hrs in a cell culture incubator. 20  $\mu$ l of serum free RPMI media containing 1  $\mu$ Ci of  $^3$ H-thymidine was added to each well for the last 6-8 hours. The cells were then harvested on 96 well GF/C filter plates and washed 5 times with water. The filters were counted in the presence of 40  $\mu$ l of scintillation fluid (Microscint 20) in a Packard Top Count counter.

#### **EXAMPLE 2**

#### Highly Purified Porcine mpl Ligand

Gel Elution Protocol

Equal amounts of affinity purified *mpl* ligand (fraction 6 eluted from the *mpl*-lgG column) and 2X Laemmli sample buffer were mixed at room temperature without reducing agent and loaded onto a Novex 4-20% polyacrylamide gel as quickly as possible. The sample was not heated. As a control, sample buffer without ligand was run in an adjacent lane. The gel was run at 4-6°C at 135 volts for approximately 2 1/4 hours. The running buffer was initially at room temperature. The gel was then removed from the gel box and the plate on one side of the gel removed.

A replica of the gel was made on nitrocellulose as follows: A piece of nitrocellulose was wet with distilled water and carefully laid on top of the exposed gel face so air bubbles were excluded. Fiducial marks were placed on the nitrocellulose and the gel plate so the replica could be accurately repositioned after staining. After approximately 2 minutes, the nitrocellulose was carefully removed, and the gel was wrapped in plastic wrap and placed in the refrigerator. The nitrocellulose was stained with Biorad's gold total protein stain by first agitating it in 3 x 10 ml 0.1% Tween 20 + 0.5 M NaCl + 0.1 M Tris-HCl pH 7.5 over approximately 45 minutes followed by 3 x 10 ml purified water over 5 minutes. The gold stain was then added and allowed to develop until the bands in the standards were visible. The replica was then rinsed with water, placed over the plastic wrap on the gel and carefully aligned with the fiducial marks. The positions of the Novex standards were marked on the gel plate and lines

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were drawn to indicate the cutting positions. The nitrocellulose and plastic wrap were then removed and the gel cut along the indicated lines with a sharp razor blade. The cuts were extended beyond the sample lanes so they could be used to determine the positions of the slices when the gel was stained. After the slices were removed, the remaining gel was silver stained and the positions of the standards and the cut marks were measured. The molecular weights corresponding to the cut positions were determined from the Novex standards.

The 12 gel slices were placed into the cells in two Biorad model 422 electroeluters. 12-14K molecular weight cutoff membrane caps were used in the cells. 50 mM ammonium bicarbonate + 0.05% SDS (approximately pH 7.8) was the elution buffer. One liter of buffer was chilled approximately 1 hour in a 4-6°C coldroom before use. Gel slices were eluted at 10 ma/cell (40 v initially) in a 4-6°C coldroom. Elution took approximately 4 hours. The cells were then carefully removed and the liquid above the frit removed with a pipet. The elution chamber was removed and any liquid above the membrane cap removed with a pipet. The liquid in the membrane cap was removed with a Pipetman and saved. Fifty  $\mu I$  aliquots of purified water were then placed in the cap, agitated and removed until all the SDS crystals dissolved. These washes were combined with the saved liquid above. Total elution sample volume was 300-500 μl per gel slice. Samples were placed in 10 mm Spectrapor 4 12-14K cutoff dialysis tubing which had been soaked several hours in purified water. They were dialyzed overnight at 4-6°C against 600 ml of phosphate buffered saline (PBS is approximately 4 mM in potassium) per 6 samples. The buffer was replaced the next morning and dialysis continued for 2.5 hours. Samples were then removed from the dialysis bags and placed in microfuge tubes. The tubes were placed on ice for 1 hour, microfuged at 14K rpm for 3 min. and the supernatants carefully removed from the precipitated SDS. The supernatants were then placed on ice for approximately 1 hour more and microfuged again for 4 min. The supernatants were diluted in phosphate buffered saline and submitted for the activity assay. Remaining samples were frozen at -70°C.

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#### EXAMPLE 3

#### Porcine mpi Ligand Microsequencing

Fraction 6 (2.6 ml) from the mpl-IgG affinity column was concentrated on a Microcon-10 (Amicon). In order to prevent the mpl ligand from absorbing to the Microcon, the membrane was rinsed with 1% SDS and 5  $\mu$ l of 10 % SDS was added to fraction 6. Sample buffer (20  $\mu$ l) of 2X was added to the fraction #6 after Microcon concentration (20  $\mu$ l) and the total volume (40  $\mu$ l) was loaded on a single lane of a 4-20 % gradient acrylamide gel (Novex). The gel was run following Novex protocol. The

gel was then equilibrated for 5 min. prior to electroblotting in 10 mM 3-(cyclohexylamino)-1-propanesulfonic acid (CAPS) buffer, pH 11.0, containing 10% methanol. Electroblotting onto Immobilon-PSQ membranes (Millipore) was carried out for 45 min. at 250 mA constant current in a BioRad Trans-Blot transfer cell (32). The PVDF membrane was stained with 0.1% Coomassie Blue R-250 in 40% methanol, 0.1% acetic acid for 1 min. and destained for 2-3 min. with 10% acetic acid in 50% methanol. The only proteins that were visible in the Mr 18,000-35,000 region of the blot had Mr of 30,000, 28,000 and 22,000.

Bands at 30, 28 and 22 kDa were subjected to protein sequencing. Automated protein sequencing was performed on a model 470A Applied Biosystem sequencer equipped with an on-line PTH analyzer. The sequencer was modified to inject 80-90% of the sample (Rodriguez, *J. Chromatogr.*, 350:217-225 [1985]). Acetone (~12 µl/l) was added to solvent A to balance the UV absorbance. Electroblotted proteins were sequenced in the Blott cartridge. Peaks were integrated with Justice Innovation software using Nelson Analytical 970 interfaces. Sequence interpretation was performed on a VAX 5900 (Henzel *et al.*, *J. Chromatogr.*, 404:41-52 [1987]). Neterminal sequences (using one letter code with uncertain residues in parenthesis) and quantity of material obtained (in brackets) is presented in Table 2'.

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TABLE 2'

		<i></i>	ipi Lige	ina Ami	no-Terminus	s Sequence	<u>s</u>
30	kDa	[1.8 pm	ol]				
	1	5	10	15	20	25	
<u> </u>	(S) P A I	P P A(C)D	PRLL	NKLLR	DD (H/S) VL	. H (G) R L	(SEQ ID NO: 30)
28	kDa	[0.5 pmol]					
ŀ	1	5	10	15	20	25	
	(S) P A F	PAXD	PRLLN	KLLR	D (H) V L (H)	GR	(SEQ ID NO: 31)
18-	22 kDa	[0.8	5 pmol]				
	1	5	10				
	XPAP	PAXDE	PRLX (N	I) (K)			(SEQ ID NO: 32)

#### **EXAMPLE 4**

#### Liquid Suspension Megakaryocytopolesis Assay

Human peripheral stem cells (PSC) (obtained from consenting patients) were diluted 5 fold with IMDM media (Gibco) and centrifuged for 15 min. at room temp. at 800 x g. The cell pellets were resuspended in IMDM and layered onto 60% Percoll (density 1.077 gm/ml) (Pharmacia) and centrifuged at 800 x g for 30 min. The

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light density mononuclear cells were aspirated at the interface and washed 2x with IMDM and plated out at 1-2 x 10<sup>6</sup> cells/ml in IMDM containing 30% FBS (1 ml final volume) in 24 well tissue culture clusters (Costar). APP or *mpl* ligand depleted APP was added to 10% and cultures were grown for 12-14 days in a humidified incubator at 37°C in 5% CO<sub>2</sub> and air. The cultures were also grown in the presence of 10% APP with 0.5 μg of *mpl*-IgG added at days 0, 2 and 4. APP was depleted of *mpl* ligand by passing APP through a *mpl*-IgG affinity column.

To quantitate megakaryocytopoiesis in these liquid suspension cultures, a modification of Solberg *et al.* was used and employs a radiolabeled murine IgG monoclonal antibody (HP1-1D) to GPIIbIIIa (provided by Dr. Nichols, Mayo Clinic). 100  $\mu$ g of HP1-1D (see Grant, B. *et al.*, Blood 69:1334-1339 [1987]). was radiolabeled with 1mCi of Na<sup>125</sup>I using Enzymobeads (Biorad, Richmond, CA) as described by the manufacturer's instructions. Radiolabeled HP1-1D was stored at -70°C in PBS containing 0.01% octyl-glucoside. Typical specific activities were 1-2 x 10<sup>6</sup> cpm/ $\mu$ g (>95% precipated by 12.5% trichloroacetic acid ).

Liquid suspension cultures were set up in triplicate for each experimental point. After 12-14 days in culture the 1ml cultures were transferred to 1.5ml eppendorf tubes and centrifuged at 800 x g for 10 min. at room temp. and the resultant cell pellets were resuspended in 100 µl of PBS containing 0.02% EDTA and 20% bovine calf serum. 10ng of 125I-HP1-1D in 50 µl of assay buffer was added to the resuspended cultures and incubated for 60 min. at room temperature (RT) with occasional shaking. Subsequently, cells were collected by centrifugation at 800 x g for 10 min. at RT and washed 2x with assay buffer. The pellets were counted for 1 min. in a gamma counter (Packard). Non-specific binding was determined by adding 1 µg of unlabeled HP1-1D for 60 min. before the addition of labeled HP1-1D. Specific binding was determined as the total 125I-HP1-1D bound minus that bound in the presence of excess unlabeled HP1-1D.

#### **EXAMPLE 5**

## Oligonucleotide PCR Primers

Based on the amino-terminal amino acid sequence obtained from the 30 kDa, 28 kDa and 18-22 kDa proteins, degenerate oligonucleotides were designed for use as polymerase chain reaction (PCR) primers (see **Table 4**). Two primer pools were synthesized, a positive sense 20 mer pool encoding amino acid residues 2-8 (*mpl* 1) and an anti-sense 21-mer pool complimentary to sequences encoding amino acids 18-24 (*mpl* 2).

PCT/US94/14553 **WO 95/18858** 

## TABLE 4 Degenerate Oligonucleotide Primer Pools

mpl 1:5' CCN GCN CCN GCN TGY GA 3' (2,048-fold degenerate)	(SEQ ID NO: 35)
mpl 2:5' NCC RTG NAR NAC RTG RTC RTC 3' (2,048-fold degenerate)	(SEQ ID NO: 36)

Porcine genomic DNA, isolated from porcine peripheral blood lymphocytes, was used as a template for PCR. The 50 µl reaction contained: 0.8 µg of porcine genomic 5 DNA in 10mM Tris-HCl (pH 8.3), 50mM KCl, 3mM MgCl<sub>2</sub>, 100 μg/ml BSA, 400 μM dNTPs, 1 µM of each primer pool and 2.5 units of Tag polymerase. Initial template denaturation was at 94°C for 8 min. followed by 35 cycles of 45 seconds at 94°C, 1 min. at 55°C and 1 min. at 72°C. The final cycle was allowed to extend for 10 min. at 10 72°C. PCR products were separated by electrophoresis on a 12% polyacrylamide gel and visualized by staining with ethidium bromide. It was reasoned that If the aminoterminal amino acid sequence was encoded by a single exon then the correct PCR product was expected to be 69 bp. A DNA fragment of this size was eluted from the gel and subcloned into pGEMT (Promega). Sequences of three clones are shown below in 15 Table 5.

#### TABLE 5

## 69 bp Porcine Genomic DNA Fragments

#### gemT3

5'CCAGCGCCGC CAGCCTGTGA CCCCCGACTC CTAAATAAAC TGCCTCGTGA 3'GGTCGCGGCG GTCGGACACT GGGGGCTGAG GATTTATTTG ACGGAGCACT

TGACCACGTT CAGCACGGC [69 bp] (SEQ ID NO: 37) ACTGGTGCAA GTCGTGCCG

(SEQ ID NO: 38)

#### gemT7

5'CCAGCACCTC CGGCATGTGA CCCCCGACTC CTAAATAAAC TGCTTCGTGA 3'GGTCGTGGAG GCCGTACACT GGGGGCTGAG GATTTATTTG ACGAAGCACT

CGACCACGTC CATCACGGC [69 bp] (SEQ ID NO: 39)

GCTGGTGCAG GTAGTGCCG

(SEQ ID NO: 40)

gemT9

PRLLNKL LR(SEQID

NO: 32)

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5' CCAGCACCGCCGGCATGTGACCCCCGACTCCTAAATAAACTGCTTCGTGACG

3' GGTCGTGGCGCCGTACACTGGGGGCTGAGGATTTATTTGACGAAGCACTGC

ATCATGTCTATCACGGT 3' (SEQ ID NO: 41)

TAGTACAGATAGTGCCA 5' (SEQ ID NO: 42)

The position of the PCR primers is indicated by the underlined bases. These results verify the N-terminal sequence obtained for amino acids 9-17 for the 30 kDa, 28 kDa and 18-22 kDa proteins and indicated that this sequence is encoded by a single exon of porcine DNA.

#### EXAMPLE 6

#### Human mpl Ligand Gene

Based on the results from **Example 5**, a 45-mer deoxyoligonucleotide, called pR45, was designed and synthesized to screen a genomic library. The 45-mer had the following sequence:

5' GCC-GTG-AAG-GAC-GTG-GTC-ACG-AAG-CAG-TTT-ATT-TAG-GAG-TCG 3' (SEQ ID NO: 28)

This oligonucleotide was  $^{32}$ P-labeled with ( $\gamma^{32}$ P)-ATP and T4 kinase and used to screen a human genomic DNA library in  $\lambda$ gem12 under low stringency hybridization and wash conditions (see **Example 7**). Positive clones were picked, plaque purified and analyzed by restriction mapping and southern blotting. Clone #4 was selected for additional analysis.

A 2.8 kb BamHI-Xbal fragment that hybridized to the 45-mer was subcloned into pBluescript SK-. Partial DNA sequencing of this clone was preformed using as primers oligonucleotides specific to the porcine *mpl* ligand DNA sequence. The sequence obtained confirmed that DNA encoding the human homolog of the porcine *mpl* ligand had been isolated. An EcoRI restriction site was detected in the sequence allowing us to isolate a 390 bp EcoRI-Xbal fragment from the 2.8 kb BamHI-Xbal and to subclone it in pBluescript SK-.

Both strands of this fragment were sequenced. The human DNA sequence and deduced amino acid sequence are shown in Fig. 9 (SEQ ID NOS: 3 & 4). The predicted positions of introns in the genomic sequence are also indicated by arrows, and define a putative exon ("exon 3").

Examination of the predicted amino acid sequence confirms that a serine residue is the first amino acid of the mature *mpl* ligand, as determined from direct amino acid sequence analysis. Immediately upstream from this codon the predicted amino acid sequence is highly suggestive of a signal sequence involved in secretion of the mature *mpl* ligand. This signal sequence coding region is probably interrupted at nucleotide position 68 by an intron.

In the 3' direction the exon appears to terminate at nucleotide 196. This exon therefore encodes a sequence of 42 amino acids, 16 of which are likely to be part of a signal sequence and 26 of which are part of the mature human *mpl* ligand.

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#### **EXAMPLE 7**

## Full Length Human mpl Ligand cDNA

Based on the human "exon 3" sequence (Example 6) two non-degenerate oligonucleotides corresponding to the 3' and 5' ends of the "exon 3" sequence were synthesized (Table 6).

TABLE 6
Human cDNA Non-degenerate PCR Oligopucleotid

Trainer Str. Non-acgenerate FCN Originacieotta P	IIIIIE19
Fwd primer: 5' GCT AGC TCT AGA AAT TGC TCC TCG TGG TCA TGC TTC T 3'	(SEQ ID NO:
	43)
Rvs primer: 5' CAG TCT GCC GTG AAG GAC ATG G 3'	(SEQ ID NO:
	44)

These two primers were used in PCR reactions employing as a template DNA from various human cDNA libraries or 1 ng of Quick Clone cDNA (Clonetech) from various tissues using the conditions described in the Example 5. The expected size of the correct PCR product was 140 bp. After analysis of the PCR products on a 12% polyacrylamide gel, a DNA fragment of the expected size was detected in cDNA libraries prepared from adult kidney, 293 fetal kidney cells and cDNA prepared from human fetal liver (Clonetech cat. #7171-1).

A fetal liver cDNA library in  $\lambda$  DR2 (Clonetech cat. # HL1151x) was screened with the same 45 mer oligonucleotide used to screen the human genomic library. The oligonucleotide was labelled with ( $\gamma^{32}$ P)-ATP using T4 polynucleotide kinase. The library was screened under low stringency hybridization conditions. The filters were prehybridized for 2hr then hybridized with the probe overnight at 42°C in 20% formamide, 5xSSC, 10xDenhardt's, 0.05M sodium phosphate (pH 6.5), 0.1% sodium pyrophosphate, 50  $\mu$ g/ml of sonicated salmon sperm DNA for 16hr. Filters were then rinsed in 2xSSC and then washed once in 0.5xSSC, 0.1% SDS at 42°C. Filters were exposed overnight to Kodak X-Ray film. Positive clones were picked, plaque purified

and the insert size was determined by PCR using oligonucleotides flanking the BamHl-Xbal cloning in  $\lambda$  DR2 (Clonetech cat. #6475-1). 5  $\mu$ I of phage stock was used as a template source. Initial denaturation was for 7 min. at 94°C followed by 30 cycles of amplification (1 min. at 94°C, 1 min. at 52°C and 1.5 min. at 72°C). Final extention was for 15 min. at 72°C. Clone # FL2b had a 1.8kb insert and was selected for further analysis.

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The plasmid pDR2 (Clonetech, \( \lambda DR2 \& pDR2 \) cloning and Expression System Library Protocol Handbook, p 42) contained within the λDR2 phage arms, was rescued as described per manufacturer's instructions (Clonetech, \( \DR2 \) & pDR2 cloning and Expression System Library Protocol Handbook, p 29-30). Restriction analysis of the plasmid pDR2-FL2b with BamHI and XbaI indicated the presence of an internal BamHI restriction site in the insert approximately at position 650. Digestion of the plasmid with BamHI-Xbal cut the insert in two fragments, one of 0.65 kb and one of 1.15 kb. DNA sequence was determined with three different classes of template derived from the plasmid pDR2-FL2b. DNA sequencing of double-stranded plasmid DNA was carried out with the ABI373 (Applied Biosystems, Foster City, California) automated fluorescent DNA sequencer using standard protocols for dye-labeled dideoxy nucleoside triphosphate terminators (dye-terminators) and custom synthesized walking primers (Sanger et al., Proc. Natl. Acad. Sci. USA, 74:5463-5467 [1977]; Smith et al., Nature, 321:674-679 [1986]). Direct sequencing of polymerase chain reaction amplified fragments from the plasmid was done with the ABI373 sequencer using custom primers and dye-terminator reactions. Single stranded template was generated with the M13 Janus vector (DNASTAR, Inc., Madison, Wisconsin) (Burland et al., Nucl. Acids Res., 21:3385-3390 [1993]). BamHl-Xbal (1.15 kb) and BamHl (0.65 kb) fragments were isolated from the plasmid pDR2-FL2b, the ends filled in with T4 DNA polymerase in the presence of deoxynucleotides, and then subcloned into the Small site of M13 Janus. Sequencing was carried out with standard protocols for dye-labeled M13 universal and reverse primers, or walking primers and dye-terminators. Manual sequencing reactions were carried out on single strand M13 DNA using walking primers and standard dideoxy-terminator chemistry (Sanger et al., Proc. Natl. Acad. Sci. USA, 74:5463-5467 [1977]),  $^{33}$ P-labeled  $\alpha$ -dATP and Sequenase (United States Biochemical Corp., Cleveland, Ohio). DNA sequence assembly was carried out with Sequencher V2.1b12 (Gene Codes Corporation, Ann Arbor, Michigan). nucleotide and deduced sequences of hML are provided in Fig. 1 (SEQ ID NO: 1).

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### **EXAMPLE 8**

# Isolation of the Human mpl Ligand (TPO) Gene.

Human genomic DNA clones of the TPO gene were isolated by screening a human genomic library in  $\lambda$ -Gem12 with pR45, a previously described oligonucleotide probe under low stringency conditions (see Example 7) or under high stringency conditions with a fragment corresponding to the 3' half of human cDNA coding for the mpl ligand (from the BamH1 site to the 3'end). Two overlapping lambda clones spanning 35 kb were isolated. Two overlapping fragments (BamH1 and EcoRI) containing the entire TPO gene were subcloned and sequenced. The structure of the human gene is composed of 6 exons within 7 kb of genomic DNA (Fig. 14 A, B and C). The boundaries of all exon/intron junctions are consistent with the consensus motif established for mammalian genes (Shapiro, M. B., et al., Nucl. Acids Res. 15:7155 [1987]). Exon 1 and exon 2 contain 5' untranslated sequence and the initial four amino acids of the signal peptide. The remainder of the secretory signal and the first 26 amino acids of the mature protein are encoded within exon 3. The entire carboxyl domain and 3' untranslated as well as ~50 amino acids of the erythropoletin-like domain are encoded within exon 6. The four amino acids involved in the deletion observed within hML-2 (hTPO-2) are encoded at the 5' end of exon 6.

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### **EXAMPLE 9**

## Transient Expression of Human mpl Ligand (hML)

In order to subclone the full length insert contained in pDR2-FL2b, the plasmid was digested with Xbal to completion, then partially digested with BamHI. A DNA fragment corresponding to the 1.8 kb insert was gel purified and subcloned in pRK5 (pRK5-hmpl I) (see U.S. Patent No. 5,258,287 for construction of pRK5) under the control of the cytomegalovirus immediate early promoter. DNA from the construct pRK5-hmpl I was prepared by the PEG method and transfected in Human embryonic kidney 293 cells maintained in Dulbecco's modified Eagle's medium (DMEM) supplemented with F-12 nutrient mixture, 20 mM Hepes (pH 7.4) and 10% fetal bovine serum. Cells were transfected by the calcium phosphate method as described (Gorman, C. [1985] in DNA Cloning: A Practical Approach (Glover, D. M., ed) Vol. II, pp. 143-190, IRL Press, Washington, D. C.). 36 h after transfection, the supernatant of the transfected cells was assayed for activity in the proliferation assay (see Example I). Supernatant of 293 cells transfected with pRK vector only gave no stimulation of the Ba/F3 or Ba/F3-mpl cells (Fig. 12A). Supernatant of cells transfected with pRK5-hmpl I had no effect on the Ba/F3 cells but dramatically stimulates the proliferation of Ba/F3-mpl cells (Fig. 12A), indicating that this cDNA encodes a functionally active human mpl ligand.

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### **EXAMPLE 10**

# Human Mpl Ligand Isoforms hML2. hML3, and hML4

In order to identify alternatively spliced forms of hML, primers were synthesized corresponding to each end of the coding sequence of hML. These primers were employed in RT-PCR to amplify human adult liver RNA. Additionally, internal primers flanking selected regions of interest (see below) were constructed and similarly employed. Direct sequencing of the ends of the PCR product revealed a single sequence corresponding exactly to the sequence of the cDNA isolated from the human fetal liver library (see Fig. 1 [SEQ ID NO: 1]). However, a region near the C-terminus of the EPO-domain (in the middle of the PCR product) exhibited a complex sequence pattern sugesting the existence of possible splice variants in that region. To isolate these splice variants, the primers provided in Table 7 flanking the region of interest were used in a PCR as templates for human adult liver cDNA.

TABLE 7

	Human ML Isoform PCR Primers	
phmpllcdna.3e1:	5'TGTGGACTTTAGCTTGGGAGAATG3'	(SEQ ID NO: 45)
pbx4.f2:		(SEQ ID NO: 46)
PDATILE	0.00.000.000	

The PCR products were subcloned blunt into M13. Sequencing of individual subclones revealed the existence of at least 3 ML isoforms. One of them, hML (also refered to as hML332), is the longest form and corresponds exactly to the sequence isolated from the fetal liver library. Sequences of the four human *mpl* ligand isoforms listed from longest (hML) to shortest (hML-4) are provided in (Fig. 11 [SEQ ID NOS: 6, 8, 9 & 10]).

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### **EXAMPLE 11**

# Construction and Transient Expression of Human Mpl Ligand Isoforms and Substitutional Variants

hML2, hML3, and hML(R153A, R154A)

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Isoforms hML2 and hML3 and substitutional variant hML(R153A, R154A) were reconstituted from hML using the recombinant PCR technique described by Russell Higuchi, in PCR Protocols, *A guide to Methods and Applications*, Acad. Press, M.A.Innis, D.H. Gelfand, J.J. Sninsky & T.J. White Editors.

In all contructs, the "outside" primers used are shown in **Table 8** and the "overlapping" primers are shown in **Table 9**.

TABLE 8
Outside Primers

CIa.FL.F2: 5'ATC GAT ATC GAT AGC CAG ACA CCC CGG CCA G3'	(SEQ ID NO:
HMPLL-R: 5'GCT AGC TCT AGA CAG GGA AGG GAG CTG TAC ATG AGA3'	(SEQ ID NO:

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TABLE 9

Overlapping Primers hML-2: ML<sub>4</sub>.F: 5'CTC CTT GGA ACC CAG GGC AGG ACC 3' (SEQ ID NO: 49) 5'GGT CCT GCC CTG GGT TCC AAG GAG 3' ML∆4.R (SEQ ID NO: 50) hML-3: hMLΔ116+: 5'CTG CTC CGA GGA AAG GAC TTC TGG ATT 3' (SEQ ID NO: 51) hMLΔ116-: 5'AAT CCA GAA GTC CTT TCC TCG GAG CAG 3' (SEQ ID NO: 52) hML(R153A, R154A): RR-KO-F: 5'CCC TCT GCG TCG CGG CGC CCC CAC CCA C 3' (SEQ ID NO: 53) RR-KO-R: 5'GTG GGT GGG GCC GCC ACG CAG AGG G 3' (SEQ ID NO: 54)

All PCR amplifications were performed with cloned Pfu DNA polymerase (Stratagene) using the following conditions: Initial template denaturation was at 94°C for 7 min. followed by 30 cycles of 1 min. at 94°C, 1 min. at 55°C and 1.5 min. at 72°C. The final cycle was allowed to extend for 10 min. at 72°C. The final PCR product was digested with Clal-Xbal, gel purified and cloned in pRK5tkneo. 293 cells were transfected with the various constructs as described above and the supernatant was assayed using the Ba/F3-mpl proliferation assay. hML-2 and hML-3 showed no detectable activity in this assay, however the activity of hML(R153A, R154A) was similar to hML indicating that processing at this di-basic site is not required for activity (see Fig. 13).

### EXAMPLE 12

# Murine mpl Ligand cDNA mML. mML-2 and mML-3

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Isolation of mML cDNA.

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A DNA fragment corresponding to the entire coding region of the human mpl ligand was obtained by PCR, gel purified and labeled by random priming in the presence of <sup>32</sup>P-dATP and <sup>32</sup>P-dCTP. This probe was used to screen 10<sup>6</sup> clones of a mouse liver cDNA library in λGT10 (Clontech cat# ML3001a). Duplicate filters were hybridized in 35% formamide, 5xSSC, 10xDenhardt's, 0.1% SDS, 0.05M sodium phosphate (pH 6.5), 0.1% sodium pyrophosphate, 100 μg/ml of sonicated salmon sperm DNA overnight in the presence of the probe. Filters were rinsed in 2xSSC and then washed once in 0.5xSSC, 0.1% SDS at 42°C. Hybridizing phage were plaque-purified and the cDNA inserts were subcloned into the Eco R1 site of Bluescript SK- plasmid. Clone "LD" with a 1.5 kb insert was chosen for further analysis and both strands were sequenced as descibed above for the human ML cDNA. The nucleotide and deduced amino acid sequences from clone LD are provided in Fig. 14 (SEQ ID NOS: 1 & 11). The deduced mature ML sequence from this clone was 331 amino acid residues long and identified as mML331 (or mML-2 for reasons described below). Considerable identity for both nucleotide and deduced amino acid sequences were observed in the EPO-like domains of these ML's. However, when deduced amino acid sequences of human and mouse ML's were aligned, the mouse sequence appeared to have a tetrapeptide deletion between human residues 111-114 corresponding to the 12 nucleotide deletion following nucleotide position 618 seen in both the human (see above) and pig (see below) cDNA's. Accordingly, additional clones were examined to detect possible murine ML isoforms. One clone, "L7", had a 1.4 kb insert with a 335 amino acid deduced sequence containing the "missing" tetrapeptide LPLQ. This form is believed to be the full length murine ML and is refered to as mML or mML335. The nucleotide and deduced amino acid sequence for mML are provided in Fig. 16 (SEQ ID NOS: 12 & 13). Finally, clone "L2" was isolated and sequenced. This clone has the 116 nucleotide deletion corresponding to hML3 and is therefore denominated mML-3. Comparison of the deduced amino acid sequences of these two isoforms is shown in Fig. 16.

Expression of recombinant mML. Expression vectors for murine ML were prepared essentially as described in Example 8. Clones encoding mML and mML-2 were subcloned into pRK5tkneo, a mammalian expression vector that provides expression under the control of the CMV promoter and an SV40 polyadenylation signal. The resulting expression vectors, mMLpRKtkneo and mML2pRKtkneo were transiently transfected into 293 cells using the calcium phosphate method. Following transient

transfection, media was conditioned for five days. The cells were maintained in high glucose DMEM media supplemented with 10% fetal calf serum.

Expression of murine-mpl (mmpl) in Ba/F3 cells. Stable cell lines expressing c-mpl were obtained by transfection of mmpl pRKtkneo, essentially as described for human mpl in Example 1. Briefly, an expression vector (20  $\mu$ g; linearized) containing the entire coding sequence of murine mpl (Skoda, R. C., et al., EMBO J. 12:2645-2653 [1993]) was transfected into Ba/F3 cells by electroporation (5 X 10<sup>6</sup> cells, 250 volts, 960  $\mu$ F) followed by selection for neomycine resistance with 2 mg/ml G418. Expression of mpl was assessed by flow cytometry analysis using rabbit anti-murine mpl-lgG antisera. Ba/F3 cells were maintained in RPMI 1640 media from WEHI -3B cells as a source of IL-3. Supernatants from 293 cells transfected with both mML and mML-2 were assayed in BaF3 cells transfected with both mmpl and hmpl as described in Example 1.

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### EXAMPLE 13

# Porcine mpl Ligand cDNA pML and pML-2

Porcine ML (pML) cDNA was isolated by RACE PCR. Briefly, an oligo dT primer and 2 specific primers were designed based on the sequence of the exon of the porcine ML gene encoding the amino terminus of the ML purified from the aplastic pig serum. cDNA prepared from various aplastic pig tissues was obtained and amplified. A PCR cDNA product of 1342 bp was found in kidney and subcloned. Several clones were sequenced and found to encode the mature pig *mpl* ligand (not including a complete secretion signal). The cDNA was found to encode a 332 amino acid mature protein (pML332) having the sequence shown in **Fig. 18** (SEQ ID NOS: 9 & 16).

#### Method:

Isolation of pML gene and cDNA. Genomic clones of the porcine ML gene were isolated by screening a pig genomic library in EMBL3 (Clontech Inc.) with pR45. The library was screened essentially as described in **Example 7**. Several clones were isolated and the exon encoding amino acid sequence identical to that obtained from the purified ML was sequenced. Porcine ML cDNA were obtained using a modification of the RACE PCR protocol. Two specific ML primers were designed based on the sequence of the pig ML gene. Polyadenylated mRNA was isolated from the kidney of aplastic pigs essentially as previously described. cDNA was prepared by reverse transcription with the BamdT primer

directed against the polyadenosine tail of the mRNA. An initial round of PCR amplification (28 cycles of 95°C for 60 seconds, 58°C for 60 seconds, and 72°C for ninety seconds) was conducted using the ML specific h-forward-1 primer

(h-forward-1: 5' GCTAGCTCTAGAAATTGCTCCTCGTGGTCATGCTTCT 3')

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(SEQ ID NO: 43)

and the BAMAD primer

(BAMAD: 5' GACTCGAGGATCCATCG 3')

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(SEQ ID NO: 56)

in a 100 ml reaction (50 mM KCl, 1.5 mM MgCl, 10 mM Tris pH 8.0, 0.2 mM dNTPs,with 0.05 U/ml Amplitaq polymerase [Perkin Elmer Inc.]) The PCR product was then digested with Cla1, extracted with phenol-chloroform (1:1), ethanol precipitated, and ligated to 0.1 mg of Bluescript SK- vector (Stratagene inc.) that had been cut with Cla1 and Kpn 1. After incubation for two hours at room temperature, one fourth of the ligation mixture was added directly to a second round of PCR (22 cycles as described above) using a second ML specific forward-1 primer

(forward-1: 5' GCTAGCTCTAGAAGCCCGGCTCCTCCTGCCTG 3')

(SEQ ID NO: 57)

and T3-21 (an oligonucleotide that binds to a sequence adjacent to the multiple cloning region within the Bluescript SK- vector):

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(5' CGAAATTAACCCTCACTAAAG 3')

(SEQ ID NO: 58).

The resulting PCR product was digested with Xba1 and Cla1 and subcloned into Bluescript SK-. Several clones from indepedent PCR reactions were sequenced.

Again, a second form, designated pML-2, encoding a protein with a 4 amino acid residue deletion (328 amino acid residues) was identified (see Fig. 21 [SEQ ID NO: 21]). Comparison of pML and pML-2 amino acid sequences shows the latter form is identical except that the tetrapeptide QLPP corresponding to residues 111-114 inclusive have been deleted (see Fig. 22 [SEQ ID NOS: 18 & 21]). The four amino acid deletions observed in murine, human and porcine ML cDNA occur at precisely the same position within the predicted proteins.

## **EXAMPLE 14**

# CMK Assay for Thrombopoietin (TPO) Induction of Platelet Antigen GPIIbIlla Expression

CMK cells are maintained in RMPI 1640 medium (Sigma) supplemented with 10% fetal bovine serum and 10mM glutamine. In preparation for the assay, the cells are harvested, washed and resuspended at 5x10<sup>5</sup> cells/ml in serum-free GIF medium supplemented with 5mg/l bovine insulin, 10mg/l apo-transferrin, 1 X trace

elements. In a 96-well flat-bottom plate, the TPO standard or experimental samples are added to each well at appropriate dilutions in 100  $\mu$ l volumes. 100  $\mu$ l of the CMK cell suspension is added to each well and the plates are incubated at 37°C, in a 5% CO2 incubator for 48 hours. After incubation, the plates are spun at 1000rpm at 4°C for five minutes. Supernatants are discarded and 100  $\mu$ l of the FITC-conjugated GPIIbIIIa monoclonal 2D2 antibody is added to each well. Following incubation at 4°C for 1 hour, plates are spun again at 1000rpm for five minutes. The supernatants containing unbound antibody are discarded and 200  $\mu$ l of 0.1% BSA-PBS wash is added to each well. The 0.1% BSA-PBS wash step is repeated three times. Cells are then analyzed on a FASCAN using standard one parameter analysis measuring relative fluorescence intensity.

### **EXAMPLE 15**

# DAMI Assay for Thrombopoietin (TPO) by Measuring Endomitotic Activity of DAMI Cells on 96-well Microtiter Plates

DAMI cells are maintained in IMDM + 10% horse serum (Gibco) supplemented with 10mM glutamine, 100ng/ml Penicillin G, and 50  $\mu$ g/ml streptomycin. In preparation for the assay, the cells are harvested, washed, and resuspended at 1x10<sup>6</sup> cells/ml in IMDM + 1% horse serum. In a 96-well round-bottom plate, 100  $\mu$ l of the TPO standard or experimental samples is added to DAMI cell suspension. Cells are then incubated for 48 hours at 37°C in a 5% CO2 incubator. After incubation, plates are spun in a Sorvall 6000B centrifuge at 1000rpm for five minutes at 4°C. Supernatants are discarded and 200  $\mu$ l of PBS-0.1% BSA wash step is repeated. Cells are fixed by the addition of 200  $\mu$ l ice-cold 70% Ethanol-PBS and resuspended by aspiration. After incubation at 4°C for 15 minutes, the plates are spun at 2000 rpm for five minutes and 150  $\mu$ l of 1mg/ml RNAse containing 0.1mg/ml propidlum iodide and 0.05% Tween-20 is added to each well. Following a one hour incubation at 37°C the changes in DNA content are measured by flow cytometry. Polyploidy is measured and quantitated as follows:

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Normalized Polyploid Ratio (NPR) = (%Cells in > G2+M/%Cells in < G2+M) with TPO (%Cells in >G2+M/%Cells in <G2+M) in control

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## EXAMPLE 16

# Thrombopoietin (TPO) In Vivo Assay (Mouse Platelet Rebound Assay)

In Vivo Assay for 35S Determination of Platelet Production

C57BL6 mice (obtained from Charles River) are injected intraperitoneally (IP) with 1 ml goat anti-mouse platelet serum (6 amps) on day 1 to produce thrombocytopenia. On days 5 and 6, mice are given two IP injections of the factor or PBS as the control. On day 7, thirty  $\mu \text{Ci}$  of Na2<sup>35</sup>SO4 in 0.1 ml saline are injected intravenously and the percent <sup>35</sup>S incorporation of the injected dose into circulating platelets is measured in blood samples obtained from treated and control mice. Platelet counts and leukocyte counts are made at the same time from blood obtained from the retro-orbital sinus.

### **EXAMPLE 17**

KIRA ELISA for Thrombopoietin (TPO)

The human mpl receptor has been disclosed by Vigon et al., PNAS, USA 89:5640-5644 (1992). A chimeric receptor comprising the extracellular domain (ECD) of the mpl receptor and the transmembrane (TM) and intracellular domain (ICD) of Rse (Mark et al., J. of Biol. Chem. 269(14):10720-10728 [1994]) with a carboxyl-terminal flag polypeptide (i.e. Rse.gD) was made for use in the KIRA ELISA described herein. See Fig. 30 and 31 for a diagrammatic description of the assay.

(a) Capture agent preparation

Monoclonal anti-gD (clone 5B6) was produced against a peptide from Herpes simplex virus glycoprotein D (Paborsky *et al.*, Protein Engineering 3(6):547-553 [1990]). The purified stock preparation was adjusted to 3.0mg/ml in phosphate buffered saline (PBS), pH 7.4 and 1.0ml aliquots were stored at -20° C.

(b) Anti-phosphotyrosine antibody preparation

Monoclonal anti-phosphotyrosine, clone 4G10, was purchased from UBI (Lake Placid, NY) and biotinylated using long-arm biotin-N-hydroxysuccinamide (Biotin-X-NHS, Research Organics, Cleveland, OH).

(c) Ligand

The *mpl* ligand was prepared by the recombinant techniques described herein. The purified *mpl* ligand was stored at 4 °C. as a stock solution.

(d) Preparation of Rse.gD nucleic acid

Synthetic double stranded oligonucleotides were used to reconstitute the coding sequence for the C-terminal 10 amino acids (880 - 890) of human Rse and add an

additional 21 amino acids containing an epitope for the antibody 5B6 and a stop codon. Table 10 presents the final sequence of the synthetic portion of the fusion gene.

#### TABLE 10

Synthetic Double Stranded Portion of Human Rse Fusion	Gene
coding strand:	
5'-TGCAGCAAGGGCTACTGCCACACTCGAGCTGCGCAGATGCTAGCCTCAAGA	
TGGCTG ATCCAAATCGATTCCGCGGCAAAGATCTTCCGGTCCTGTAGAAGCT-3'	(SEQ ID
	NO: 59)
noncoding (anti-sense) strand:	
5'-AGCTTCTACAGGACCGGAAGATCTTTGCCGCGGAATCGATTTGGATCAGCCA	
TCTTG AGGCTAGCATCTGCGCAGCTCGAGTGTGGCAGTAGCCCTTGCTGCA-3'	(SEQ ID
	NO: 60)

The synthetic DNA was ligated with the cDNA encoding amino acids 1-880 of human Rse at the Pstl site beginning at nucleotide 2644 of the published human Rse cDNA sequence (Mark et al., Journal of Biological Chemistry 269(14):10720-10728 [1994]) and HindIII sites in the polylinker of the expression vector pSVI7.ID.LL (See Fig. 32 A-L; SEQ ID NO: 22) to create the expression plasmid pSV.ID.Rse.gD. Briefly, the expression plasmid comprises a dicistronic primary transcript which contains sequence encoding DHFR bounded by 5' splice donor and 3' splice acceptor intron splice sites, followed by sequence that encodes the Rse.gD. The full length (non-spliced) message contains DHFR as the first open reading frame and therefore generates DHFR protein to allow selection of stable transformants.

### (e) Preparation of mpl-Rse.gD nucleic acid

The expression plasmid pSV.ID.Rse.gD produced as described above was modified to produce plasmid pSV.ID.M.tmRd6 which contained the coding sequences of the ECD of human *mpl* (amino acids 1-491) fused to the transmembrane domain and intracellular domain of Rse.gD (amino acids 429-911). Synthetic oligonucleotides were used to join the coding sequence of a portion of the extracellular domain of human *mpl* to a portion of the Rse coding sequence in a two step PCR cloning reaction as described by Mark *et al.*, *J. Biol. Chem.* 267:26166-26171 (1992). Primers used for the first PCR reaction were M1

(5'-TCTCGCTACCGTTTACAG-3')

(SEQ ID NO: 61)

and M2

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# (5'-CAGGTACCCACCAGGCGGTCTCGGT-3')

(SEQ ID NO: 62)

with a mpl cDNA template and R1

(5'-GGGCCATGACACTGTCAA-3')

(SEQ ID NO: 63)

and R2

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(5'-GACCGCCACCGAGACCGCCTGGTGGGTACCTGTGGTCCTT-3')

(SEQ ID NO: 64)

with a Rse cDNA template. The Pvull-Smal portion of this fusion junction was used for the construction of the full-length chimeric receptor.

### (f) Cell transformation

DP12.CHO cells (EP 307,247 published 15 March 1989) were electroporated with pSV.ID.M.tmRd6 which had been linearized at a unique NotI site in the plasmid backbone. The DNA was ethanol precipitated after phenol/chloroform extraction and was resuspended in 20μl 1/10 Tris EDTA. Then, 10μg of DNA was incubated with 10<sup>7</sup> CHO DP12 cells in 1 ml of PBS on ice for 10 min. before electroporation at 400 volts and 330μf. Cells were returned to ice for 10 min. before being plated into non-selective medium. After 24 hours cells were fed nucleoside-free medium to select for stable DHFR+ clones.

(g) Selection of transformed cells for use in the KIRA ELISA

Clones expressing MPL/Rse.gD were identified by western-blotting of whole cell lysates post-fractionation by SDS-PAGE using the antibody 5B6 which detects the gD epitope tag.

## (h) Media

Cells were grown in F12/DMEM 50:50 (Gibco/BRL, Life Technologies, Grand Island, NY). The media was supplemented with 10% diafiltered FBS (HyClone, Logan, Utah), 25mM HEPES and 2mM L-glutamine.

### (i) KIRA ELISA

Mpl-Rse.gD transformed DP12.CHO cells were seeded (3x10<sup>4</sup> per well) in the wells of a flat-bottom-96 well culture plate in 100 μl media and cultured overnight at 37 °C in 5% CO<sub>2</sub>. The following morning the well supernatants were decanted, and the plates were lightly tamped on a paper towel. 50μl of media containing either experimental samples or 200, 50, 12.5, 3.12, 0.78, 0.19, 0.048 or 0 ng/ml mpl ligand was then added to each well. The cells were stimulated at 37°C for 30 min., the well supernatants were decanted, and the plates were once again lightly tamped on a paper towel. To lyse the cells and solubilize the chimeric receptors, 100 μl of lysis buffer was added to each well. Lysis buffer consisted of 150 mM NaCl containing 50 mM HEPES (Gibco), 0.5 % Triton-X 100 (Gibco), 0.01 % thimerosal, 30 KIU/ml

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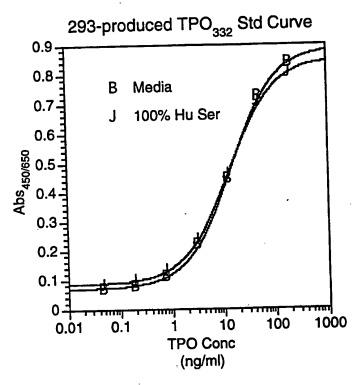
aprotinin (ICN Biochemicals, Aurora, OH), 1mM 4-(2-aminoethyl)-benzenesulfonyl fluoride hydrochloride (AEBSF; ICN Biochemicals), 50  $\mu$ M leupeptin (ICN Biochemicals), and 2 mM sodium orthovanadate (Na<sub>3</sub>VO<sub>4</sub>; Sigma Chemical Co, St. Louis, MO), pH 7.5. The plate was then agitated gently on a plate shaker (Bellco Instruments, Vineland, NJ) for 60 min. at room temperature.

While the cells were being solubilized, an ELISA microtiter plate (Nunc Maxisorp, Inter Med, Denmark) coated overnight at 4°C with the 5B6 monoclonal antigD antibody (5.0 μg/ml in 50 mM carbonate buffer, pH 9.6, 100 μl/well) was decanted, tamped on a paper towel and blocked with 150 μl/well of Block Buffer [PBS containing 0.5 % BSA (Intergen Company, Purchase, NY) and 0.01 % thimerosal] for 60 min. at room temperature with gentle agitation. After 60 minutes, the anti-gD 5B6 coated plate was washed 6 times with wash buffer (PBS containing 0.05 % Tween-20 and 0.01 % thimerosal) using an automated plate washer (ScanWasher 300, Skatron Instruments, Inc, Sterling, VA).

The lysate containing solubilized MPL/Rse.gD from the cell-culture microtiter well was transferred (85 μl/well) to anti-gD 5B6 coated and blocked ELISA well and was incubated for 2 h at room temperature with gentle agitation. The unbound mpl-Rse.gD was removed by washing with wash buffer and 100 µl of biotinylated 4G10 (anti-phosphotyrosine) diluted 1:18000 in dilution buffer (PBS containing 0.5 % BSA, 0.05 % Tween-20, 5 mM EDTA, and 0.01 % thimerosal), i.e. 56 ng/ml was added to each well. After incubation for 2 hr at room temperature the plate was washed and 100 µl of horseradish peroxidase (HRPO)-conjugated streptavidin (Zymed Laboratories, S. San Francisco, CA) diluted 1:60000 in dilution buffer was added to each well. The plate was incubated for 30 minutes at room temperature with gentle agitation. The free avidin-conjugate was washed away and 100 µl freshly prepared substrate solution (tetramethyl benzidine [TMB]; 2-component substrate kit; Kirkegaard and Perry, Gaithersburg, MD) was added to each well. The reaction was allowed to proceed for 10 minutes, after which the color development was stopped by the addition of 100 µl/well 1.0 M H<sub>3</sub>PO<sub>4</sub>. The absorbance at 450 nm was read with a reference wavelength of 650 nm (ABS450/650), using a vmax plate reader (Molecular Devices, Palo Alto, CA) controlled with a Macintosh Centris 650 (Apple Computers, Cupertino, CA) and DeltaSoft software (BioMetallics, Inc. Princeton, NJ).

The standard curve was generated by stimulating dp12.trkA,B or C.gD cells with 200, 50, 12.5, 3.12, 0.78, 0.19, 0.048 or 0 ng/ml *mpl* ligand and presented as ng/ml TPO vs. mean ABS450/650 ± sd using the DeltaSoft program. Sample concentrations were obtained by interpolation of their absorbance on the standard curve and are expressed in terms of ng/ml TPO activity.

The *mpl*-ligand was found to be able to activate the *mpl*-Rse.gD chimeric receptor in a concentration-dependent and ligand-specific manner. Further, the *mpl*-Rse.gD KIRA-ELISA was found to be tolerant of up to 100% human serum (shown) or 100% plasma (not shown), allowing the assay to be used to readily screen patient and pK samples.



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Summary	of TPO EC50"	S
TPO Form (cells)	EC50 (wt/vol)	EC50 (molarity)
Hu TPO 332 (293)	2.56 ng/ml	67.4 pM
Mu TPO 332 (293)	3.69 ng/ml	97.1 pM
Hu TPO 153 (293)	~41 ng/ml	~1.08 nM
Hu TPO 155 (E. coli )	0.44 ng/ml	11.6 pM
Hu TPO 153met (E. coli)	0.829 ng/ml	21.8 pM

### EXAMPLE 18

### Receptor Based ELISA for Thrombopoletin (TPO)

ELISA plates were coated with rabbit F(ab')2 anti-human IgG (Fc) in pH 9.6 carbonate buffer at 4°C overnight. Plates were blocked with 0.5 % bovine serum albumin in PBS at room temperature for one hour. Fermenter harvest containing the chimeric receptor, *mpl*-IgG, was added to the plates and incubated for 2 hours. Twofold serial dilutions (0.39-25 ng/ml) of the standard (TPO332 produced in 293 cells with the concentration determined by quantitative amino acid analysis) and serially diluted samples in 0.5% bovine serum albumin, 0.05% tween 20 were added to the plates and incubated for 2 hours. Bound TPO was detected with protein A purified, biotinylated rabbit antibodies to TPO155 which was produced in E. coli (1 hour incubation), followed by streptavidin-peroxidase (30 min. incubation) and 3,3',5,5'-tetramethyl benzidine as the substrate. The absorbance was read at 450 nm. Plates were washed between steps. For data analysis, the standard curve is fitted using a four-parameter curve fitting program by Kaleidagraph. Concentrations of the samples were calculated from the standard curve.

### **EXAMPLE 19**

#### Expression and Purification of TPO from 293 Cells

### 20 1. Preparation of 293 Cell Expression Vectors

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A cDNA corresponding to the TPO entire open reading frame was obtained by PCR using the following oligonucleotides as primers:

# TABLE 11

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Cla.FL.F: 5' ATC GAT ATC GAT CAG CCA GAC ACC CCG GCC AG 3'	(SEQ ID NO: 65)
hmpli-R: 5' GCT AGC TCT AGA CAG GGA AGG GAG CTG TAC ATG AGA 3'	(SEQ ID NO:
	48)

PRK5-hmpl I (described in Example 9) was used as template for the reaction in the presence of pfu DNA polymerase (Stratagene). Initial denaturation was for 7 min. at 94°C followed by 25 cycles of amplification (1 min. at 94°C, 1 min. at 55°C and 1 min. at 72°C). Final extension was for 15 min. at 72°C). The PCR product was purified and cloned between the restriction sites Clal and Xbal of the plasmid pRK5tkneo, a pRK5 derived vector modified to express a neomycin resistance gene under the control of the thymidine kinase promote, to obtain the vector pRK5tkneo.ORF. A second construct corresponding to the epo homologous domain was

generated the same way but using Cla.FL.F as forward primer and the following reverse primer:

Arg.STOP.Xba: 5' TCT AGA TCT AGA TCA CCT GAC GCA GAG GGT GGA CC 3' (SEQ ID NO: 66)

The final construct is called pRK5-tkneoEPO-D. The sequence of both constructs was verified as described in Example 7.

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# 2. Transfection of Human Embryonic Kidney cells

These 2 constructs were transfected into Human Embryonic Kidney cells by the CaPO<sub>4</sub> method as described in **Example 9**. 24 hours after transfection selection of neomycin resistant clones was started in the presence of 0.4 mg/ml G418.10 to 15 days later individual colonies were transferred to 96 well plates and allowed to grow to confluency. Expression of ML153 or ML332 in the conditioned media from these clones was assessed using the Ba/F3-mpl proliferation assay (described in **Example 1**).

## 15 3. Purification of rhML332

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(pharmacia) column that was equilibrated in 10mM sodium phosphate pH 7.4 (buffer A). The column was subsequently washed with 10 column volumes each of buffer A and buffer A containing 2M urea. The column was then eluted with buffer A containing 2M urea and 1M NaCl. The Blue-Sepharose elution pool was then directly applied to a WGA-Sepharose column equilibrated in buffer A. The WGA-Sepharose column was then washed with 10 column volumes of buffer A containing 2M urea and 1 M NaCl and eluted with the same buffer containing 0.5M N-acetyl-D-glucosamine. The WGA-Sepharose eluate was applied to a C4-HPLC column (Synchrom, Inc.) equilibrated in 0.1% TFA. The C4-HPLC column was eluted with discontinuous propanol gradient (0-25%, 25-35%, 35-70%). rhML332 was found to elute in the 28-30% propanol region of the gradient. By SDS-PAGE the purified rhML332 migrates as a broad band in the 68-80 kDa region of the gel(see Figure 15).

## 4. Purification of rhML<sub>153</sub>

293-rhML<sub>153</sub> conditioned media was resolved on Blue-Sepharose as described for rhML<sub>332</sub>. The Blue Sepharose eluate was applied directly to a *mpl*-affinity column as described above. RhML<sub>153</sub> eluted from the *mpl*-affinity column was purified to homogeneity using a C4-HPLC column run under the same conditions as described for rhML<sub>332</sub>. By SDS-PAGE the purified rhML<sub>153</sub> resolves into 2 major and 2 minor bands with Mr of \_18,000-21,000(see **Figure 15**).

### **EXAMPLE 20**

#### Expression and Purification of TPO from CHO

1. Description of CHO Expression Vectors

The expression vectors used in the electroporation protocols described below based below have been designated:

pSVI5.ID.LL.MLORF (full length or hTPO332), and pSVI5.ID.LL.MLEPO-D (truncated or hTPO153).

The pertinent features of these plasmids are presented in Fig. 23 and 24.

2. Preparation of CHO Expression Vectors

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A cDNA corresponding to the hTPO entire open reading frame was obtained by PCR using the oligonucleotide primers of **Table 12**.

TABLE 12
CHO Expression Vector PCR Primers

Cla.FL.F2	5' ATC GAT ATC GAT AGC CAG ACA CCC CGG CCA G 3'	(SEQ ID NO:
ORF. Sal	5' AGT CGA CGT CGA CGT CGG CAG TGT CTG AGA ACC 3'	(SEQ ID NO:
		67)

PRK5-hmpl I (described in Example 7 and 9) was used as template for the reaction in the presence of pfu DNA polymerase (Stratagene). Initial denaturation was for 7 min. at 94°C followed by 25 cycles of amplification (1 min. at 94°C, 1 min. at 55°C and 1 min. at 72°C). Final extension was for 15 min. at 72°C). The PCR product was purified and cloned between the restriction sites Clal and Sall of the plasmid pSVI5.ID.LL to obtain the vector pSVI5.ID.LL.MLORF. A second construct corresponding to the EPO homologous domain was generated the same way but using Cla.FL.F2 as forward primer and the following reverse primer:

EPOD.Sal 5' AGT CGA CGT CGA CTC ACC TGA CGC AGA GGG TGG ACC 3' (SEQ ID NO: 68)

The final construct is called pSVI5.ID.LL.MLEPO-D. The sequence of both constructs was verified as described in **Example 7** and **9**.

in essence, the coding sequences for the full length and truncated ligand were introduced into the multiple cloning site of the CHO expression vector pSVI5.ID.LL. This vector contains the SV40 early promoter/enhancer region, a modified splice unit containing the mouse DHFR cDNA, a multiple cloning site for the introduction of the gene of interest (in this case the TPO sequences described) an SV40 polyadenylation signal and origin of replication and the beta-lactamase gene for plasmid selection and amplification in bacteria.

3. Methodology for Establishing Stable CHO Cell Lines Expressing Recombinant Human TPO332 and TPO153

# a. Description of CHO parent cell line

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The host CHO (Chinese Hamster Ovary) cell line used for the expression of the TPO molecules described herein is known as CHO-DP12 (see EP 307,247 published 15 March 1989). This mammalian cell line was clonally selected from a transfection of the parent line (CHO-K1 DUX-B11(DHFR-)- obtained from Dr. Frank Lee of Stanford University with the permission of Dr.L. Chasin) with a vector expressing preproinsulin to obtain clones with reduced insulin requirements. These cells are also DHFR minus and clones can be selected for the presence of DHFR cDNA vector sequences by growth on medium devoid of nucleoside supplements (glycine, hypoxanthine, and thymidine). This selection system for stably expressing CHO cell lines is commonly used.

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# b. Transfection method (electroporation)

TPO332 and TPO153 expressing cell lines were generated by transfecting DP12 cells via electroporation (see e.g. Andreason, G.L. J. Tiss. Cult. Meth., 15,56 [1993]) with linearized pSVI5.ID.LL.MLORF or pSVI5.ID.LL.MLEPO-D plasmids respectively. Three (3) restriction enzyme reaction mixtures were set up for each plasmid cutting; 10μg, 25μg and 50μg of the vector with the enzyme NOTI by standard molecular biology methods. This restriction site is found only once in the vector in the linearization region 3' and outside the TPO ligand transcription units (see Fig. 23). The 100μl reactions were set up for overnight incubation at 37 degrees. The next day the mixes were phenol-chloroform-isoamyl alcohol (50:49:1) extracted one time and ethanol precipitated on dry ice for approximately one hour. The precipitate was then collected by a 15 minute microcentrifugation and dried. The linearized DNA was resuspended into 50μl of Ham's DMEM-F12 1:1 medium supplemented with standard antibiotics and 2mM glutamine.

Suspension growing DP12 cells were collected, washed one time in the medium described for resuspending the DNA and finally resuspended in the same medium at a concentration of 10<sup>7</sup> cells per 750µl. Aliquots of cells (750µl) and each linearized DNA mix were incubated together at room temperature for one hour and then transferred to a BRL electroporation chamber. Each reaction mix was then electroporated in a standard BRL electroporation apparatus at 350 volts set at 330µF and low capacitance. After electroporation, the cells were allowed to sit in the apparatus for 5 minutes and then on ice for an additional 10 minute incubation period. The electroporated cells were transferred to 60mm cell culture dishes containing 5 ml of standard, complete growth medium for CHO cells (High glucose DMEM-F12 50:50

without glycine supplemented with 1X GHT, 2mM glutamine, and 5% fetal calf serum) and grown overnight in a 5% CO<sub>2</sub> cell culture incubator.

### c. Selection and screening method

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The next day, cells were trypsinized off the plates by standard methods and transferred to 150mm tissue culture dishes containing DHFR selective medium (Ham's DMEM-F12, 1:1 medium described above supplemented with either 2% or 5% dialyzed fetal calf serum but devoid of glycine, hypoxanthine and thymidine this is the standard DHFR selection medium we use). Cells from each 60mm dish were subsequently replated into 5 /150 mm dishes. Cells were then incubated for 10 to 15 days( with one medium change) at 37 degrees/5% CO2 until clones began to appear and reached sizes amenable to transfer to 96 well dishes. Over a period of 4-5 days, cell lines were transferred to 96 well dishes using sterile yellow tips on a pipettman set at 50ml. The cells were allowed to grow to confluency (usually 3-5 days) and then the trays were trypsinized and 2 copies of the original tray were reproduced. Two of these copies were short term stored in the freezer with cells in each well diluted into 50ul of 10%FCS in DMSO. 5 day conditioned serum free medium samples were assayed from confluent wells in the third tray for TPO expression via the Ba/F cell based activity assay. The highest expressing clones based on this assay were revived from storage and scaled up to 2 confluent 150mm T-flasks for transfer to the cell culture group for suspension adaptation, re-assay and banking.

### d. Amplification Protocol

Several of the highest titer cell lines from the selection described above were subsequently put through a standard methotrexate amplification regime to generate higher titer clones. CHO cell clones are expanded and plated in 10cm dishes at 4 concentrations of methotrexate (*i.e.*. 50nM, 100nM, 200nM and 400nM) at two or three cell numbers (105, 5x105, and 106 cells per dish). These cultures are then incubated at 37 degree/5% CO<sub>2</sub> until clones are established and amenable to transfer to 96 well dishes for further assay. Several high titer clones from this selection were again subjected to greater concentrations of methotrexate (*i.e.* 600nM, 800 nM, 1000nM and 1200nM) and as before resistant clones are allowed to establish and then transferred to 96 well dishes and assayed.

# 4. Culturing Stable CHO Cell Lines Expressing Recombinant Human TPO332 and TPO153

Banked cells are thawed and the cell population is expanded by standard cell growth methods in either serum free or serum containing medium. After expansion to sufficient cell density, cells are washed to remove spent cell culture media. Cells are then cultured by any standard method including; batch, fed-batch or continuous culture at 25-40 °C, neutral pH, with a dissolved O<sub>2</sub> content of at least 5% until the

constitutively secreted TPO is accumulated. Cell culture fluid is then separated from the cells by mechanical means such as centrifugation.

5 Purification of Recombinant Human TPO from CHO Culture Fluids

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Harvested cell culture fluid (HCCF) is directly applied to a Blue Sepharose 6 Fast Flow column (Phamacia) equilibrated in 0.01M Na Phosphate pH7.4, 0.15M NaCl at a ratio of approximately 100L of HCCF per liter of resin and at a linear flow rate of approximately 300 ml/hr/cm<sup>2</sup>. The column is then washed with 3 to 5 column volumes of equilibration buffer followed by 3 to 5 column volumes of 0.01M Na Phosphate pH7.4, 2.0M urea. The TPO is then eluted with 3 to 5 column volumes of 0.01M Na Phosphate pH7.4, 2.0M urea, 1.0M NaCl.

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The Blue Sepharose Pool containing TPO is then applied to a Wheat Germ Lectin Sepharose 6MB column (Pharmacia) equilibrated in 0.01M Na Phosphate pH7.4, 2.0M urea, and 1.0M NaCl at a ratio of from 8 to 16 ml of Blue Sepharose Pool per ml of resin at flow rate of approximately 50 ml/hr/cm<sup>2</sup>. The column is then washed with 2 to 3 column volumes of equilibration buffer. The TPO is then eluted with 2 to 5 column volumes of 0.01M Na Phosphate pH7.4, 2.0M urea, 0.5M N-acetyl-D-glucosamine.

The Wheat Germ Lectin Pool is then adjusted to a final concentration of 0.04% C12E8 and 0.1% trifluroacetic acid (TFA). The resulting pool is applied to a C4 reverse phase column (Vydac 214TP1022) equilibrated in 0.1% TFA, 0.04% C12E8 at a load of approximately 0.2 to 0.5 mg protein per ml of resin at a flow rate of 157 ml/hr/cm<sup>2</sup>.

The protein is eluted in a two phase linear gradient of acetonitrile containing 0.1% TFA, 0.04% C12E8. The first phase is composed of a linear gradient from 0 to 30% acetonitrile in 15 minutes, The second phase is composed of a linear gradient from 30 to 60% acetonitrile in 60 minutes. The TPO elutes at approximately 50% acetonitrile. A pool is made on the basis of SDS-PAGE.

The C4 Pool is then diluted with 2 volumes of 0.01M Na Phosphate pH7.4, 0.15M NaCl and diafilitered versus approximately 6 volumes of 0.01M Na Phosphate pH7.4, 0.15M NaCl on an Amicon YM or like ultrafiltration membrane having a 10,000 to 30,000 Dalton molecular weight cut-off. The resulting diafiltrate may be then directly processed or further concentrated by ultrafiltration. The diafiltrate/concentrate is adjusted to a final concentration of 0.01% Tween-80.

All or a portion of the diafiltrate/concentrate equivalent to 2 to 5% of the calculated column volume is then applied to a Sephacryl S-300 HR column (Pharmacia) equilibrated in 0.01M Na Phosphate pH7.4, 0.15M NaCl, 0.01% Tween-80 and chromatographed at a flow rate of approximately 17 ml/hr/cm<sup>2</sup>. The TPO containing fractions which are free of aggregate and proteolytic degradation products

are pooled on the basis of SDS-PAGE. The resulting pool is filtered on a  $0.22\mu$  filter, Millex-GV or like, and stored at 2-8°C.

#### **EXAMPLE 21**

Transformation and Induction of TPO Protein Synthesis in E. coli

1. Construction of E. coli TPO expression vectors

The plasmids pMP21, pMP151, pMP41, pMP57 and pMP202 are all designed to express the first 155 amino acids of TPO downstream of a small leader which varies among the different constructs. The leaders provide primarily for high level translation initiation and rapid purification. The plasmids pMP210-1, -T8, -21, -22, -24, -25 are designed to express the first 153 amino acids of TPO downstream of an initiation methionine and differ only in the codon usage for the first 6 amino acids of TPO, while the plasmid pMP251 is a derivative of pMP210-1 in which the carboxy terminal end of TPO is extended by two amino acids. All of the above plasmids will produce high levels of intracellular expression of TPO in E. coli upon induction of the tryptophan promoter (Yansura, D. G. et. al. Methods in Enzymology (Goeddel, D. V., Ed.) 185:54-60, Academic Press, San Diego [1990]). The plasmids pMP1 and pMP172 are intermediates in the construction of the above TPO intracellular expression plasmids.

20 (a) Plasmid pMP1

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The plasmid pMP1 is a secretion vector for the first 155 amino acids of TPO, and was constructed by ligating together 5 fragments of DNA as shown in Fig. 33. The first of these was the vector pPho21 in which the small Mlul-BamHI fragment had been removed. pPho21 is a derivative of phGH1 (Chang, C. N. et. al., Gene 55:189-196 [1987]) in which the human growth hormone gene has been replaced with the E. coli phoA gene, and a Mlul restriction site has been engineered into the coding sequence for the STII signal sequence at amino acids 20-21.

The next two fragments, a 258 base pair Hinfi-Psti piece of DNA from pRK5-hmpl I (Example 9) encoding TPO amino acids 19-103, and the following synthetic DNA encoding amino acids 1-18

5'-CGCGTATGCCAGCCCGGCTCCTCCTGCTTGTGACCTCCGAGTCCTCAGTAAACTGCTTCG
TG

ATACGGTCGGGCCGAGGAGGACGAACACTGGAGGCTCAGGAGTCATTTGACGAAGC
3 5 ACTGA-5'

(SEQ ID NO: 69) (SEQ ID NO: 70)

were preligated with T4-DNA ligase, and second cut with Pstl. The fourth was a 152 base pair Pstl-HaellI fragment from pRK5hmpll encoding amino acids 104-155 of TPO. The last was a 412 base pair Stul-BamHI fragment from pdh108 containing the lambda to transcriptional terminator as previously described (Scholtissek, S. et. al., NAR 15:3185 [1987]).

# (b) Plasmid pMP21

The plasmid pMP21 is designed to express the first 155 amino acids of TPO with the aid of a 13 amino acid leader comprising part of the STII signal sequence. It was constructed by ligating together three (3) DNA fragments as shown in Fig. 34, the first of these being the vector pVEG31 in which the small Xbal-SphI fragment had been removed. The vector pVEG31 is a derivative of pHGH207-1 (de Boer, H. A. et. al., in Promoter Structure and Function (Rodriguez, R. L. and Chamberlain, M. J., Ed), 462, Praeger, New York [1982]) in which the human growth hormone gene has been replaced by the gene for vascular endothelial growth factor (this identical vector fragment can be obtained from this latter plasmid).

The second part in the ligation was a synthetic DNA duplex with the following sequence:

# 5'-CTAGAATTATGAAAAAGAATATCGCATTTCTTAA

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TTAATACTTTTTCTTATAGCGTAAAGAAGAATTGCGC-5'

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(SEQ ID NO: 71)

(SEQ ID NO: 72)

The last piece was a 1072 base pair Mlul-Sphl fragment from pMP1 encoding 155 amino acids of TPO.

## (c) Plasmid pMP151

The plasmid pMP151 is designed to express the first 155 amino acids of TPO downstream of a leader comprising 7 amino acids of the STII signal sequence, 8 histidines, and a factor Xa cleavage site. As shown in Fig. 35, pMP151 was constructed by ligating together three DNA fragments, the first of these being the previously described vector pVEG31 from which the small Xbal-Sphi fragment had been removed. The second was a synthetic DNA duplex with the following sequence:

5'-CTAGAATTATGAAAAAGAATATCGCATTTCATCACCATCACCATCACCATCACATCGAAG GTCGTAGCC

TTAATACTTTTTCTTATAGCGTAAAGTAGTGGTAGTGGTAGTGGTAGCTTC
CAGCAT-5'

(SEQ ID NO: 73)

(SEQ ID NO: 74)

The last was a 1064 base pair Bgll-Sphl fragment from pMP11 encoding 154 amino acids of TPO. The plasmid pMP11 is identical to pMP1 with the exception of a few codon changes in the STII signal sequence (this fragment can be obtained from pMP1).

### (d) Plasmid pMP202

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The plasmid pMP202 is very similar to the expression vector pMP151 with the exception that the factor Xa cleavage site in the leader has been replaced with a thrombin cleavage site. As shown in Fig. 36, pMP202 was constructed by ligating together three DNA fragments. The first of these was the previously described pVEG31 in which the small Xbal-SphI fragment had been removed. The second was a synthetic DNA duplex with the following sequence:

5'-CTAGAATTATGAAAAAGAATATCGCATTTCATCACCATCACCATCACCATCACCATCGAA CCACGTAGCC

TTAATACTTTTCTTATAGCGTAAAGTAGTGGTAGTGGTAGTGTAGCTT GGTGCAT-5'

(SEQ ID NO: 75)

(SEQ ID NO: 76)

The last piece was a 1064 base pair Bgll-Sphl fragment from the previously described plasmid pMP11.

### 20 (e) Plasmid pMP172

The plasmid pMP172 is a secretion vector for the first 153 amino acids of TPO, and is an intermediate for the construction of pMP210. As shown in Fig. 37, pMP172 was prepared by ligating together three DNA fragments, the first of which was the vector pLS32lamB in which the small EcoRI-HindIII section had been removed. The second was a 946 base pair EcoRI-Hgal fragment from the previously described plasmid pMP11. The last piece was a synthetic DNA duplex with the following sequence:

5'-TCCACCCTCTGCGTCAGGT

(SEQ ID NO: 77)

GGAGACGCAGTCCATCGA-5'

(SEQ ID NO: 78)

### 30 (f) Plasmid pMP210

The plasmid pMP210 is designed to express the first 153 amino acids of TPO after a translational initiation methionine. This plasmid was actually made as a bank of plasmids in which the first 6 codons of TPO were randomized in the third position of each codon, and was constructed as shown in Fig. 38 by the ligation of three DNA fragments. The first of these was the previously described vector pVEG31 in which the small Xbal-Sphl fragment had been removed. The second was a synthetic DNA dupex shown below treated first with DNA polymerasel (Klenow) followed by digestion with

Xbal and Hinfl, and encoding the initation methionine and the randomized first 6 codons of TPO.

5'-GCAGCAGTTCTAGAATTATGTCNCCNGCNCCNCCNGCNTGTGACCTCCGA ACACTGGAGGCT

GTTCTCAGTAAA (SEQ ID NO: 79)

CAAGAGTCATTTGACGAAGCACTGAGGGTACAGGAAG-5' (SEQ ID NO: 80)

The third was a 890 base pair Hinfl-SphI fragment from pMP172 encoding amino acids 19-153 of TPO.

The plasmid pMP210 bank of approximately 3700 clones was retransformed onto high tetracycline (50  $\mu$ g/ml) LB plates to select out high translational initiation clones (Yansura, D. G. et. al., Methods: A Companion to Methods in Enzymology 4:151-158 [1992]). Of the 8 colonies which came up on high tetracycline plates, five of the best in terms of TPO expression were subject to DNA sequencing and the results are shown in Fig. 39 (SEQ ID NOS: 23, 24, 25, 26, 27 and 28).

# (g) Plasmid pMP41

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The plasmid pMP41 is designed to express the first 155 amino acids of TPO fused to a leader consisting of 7 amino acids of the STII signal sequence followed by a factor Xa cleavage site. The plasmid was constructed as shown in **Fig. 40** by ligating together three pieces of DNA, the first of which was the previously described vector pVEG31 in which the small Xbal-SphI fragment had been removed. The second was the following synthetic DNA duplex:

5'-CTAGAATTATGAAAAAGAATATCGCATTTATCGAAGGTCGTAGCC (SEQ ID NO: 81)

TTAATACTTTTCTTATAGCGTAAATAGCTTCCAGCAT-5' (SEQ ID NO: 82)

The last piece of the ligation was the 1064 base pair Bgll-Sphl fragment from the previously described plasmid pMP11.

### (h) Plasmid pMP57

The plasmid pMP57 expresses the first 155 amino acids of TPO downstream of a leader consisting of 9 amino acids of the STII signal sequence and the dibasic site Lys-Arg. This dibasic site provides for a means of removing the leader with the protease ArgC. This plasmid was constructed as shown in Fig. 41 by ligating together three DNA pieces. The first of these was the previously described vector pVEG31 in which the small Xbal-SphI fragment had been removed. The second was the following synthetic DNA duplex:

5'-CTAGAATTATGAAAAAGAATATCGCATTTCTTTAAACGTAGCC (SEQ ID NO: 83)
TTAATACTTTTTCTTATAGCGTAAAGAAGAATTTGCAT-5' (SEQ ID NO: 84)

The last part of the ligation was the 1064 base pair Bgll-Sphl fragment from the previously described plasmid pMP11.

### (i) Plasmid pMP251

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The plasmid pMP251 is a derivative of pMP210-1 in which two additional amino acids of TPO are included on the carboxy terminal end. As shown in Fig.42, this plasmid was constructed by ligating together two pieces of DNA, the first of these being the previously described pMP21 in which the small Xbal-Apal fragment had been removed. The second part of the ligation was a 316 base pair Xbal-Apal fragment from pMP210-1.

10 2. Transformation and Induction of E. coli with TPO expression vectors

The above TPO expression plasmids were used to transform the *E. coli* strain 44C6 (w3110 tonA $_{\Delta}$  rpoHts lon $_{\Delta}$  clpP $_{\Delta}$  galE) using the CaCl2 heat shock method (Mandel, M. *et al.*, *J. Mol. Biol.*, 53:159-162, [1970]). The transformed cells were grown first at 37°C in LB media containing 50 µg/ml carbenicillin until the optical density (600nm) of the culture reached approximately 2-3. The LB culture was then diluted 20x into M9 media containing 0.49% casamino acids (w/v) and 50 µg/ml carbenicillin. After growth with aeration at 30°C for 1 hour, indole-3-acrylic acid was added to a final concentration of 50 µg/ml. The culture was then allowed to continue growing at 30°C with aeration for another 15 hours at which time the cells were harvested by centrifugation.

### **EXAMPLE 22**

Production of Biologically Active TPO (Met-11-153) in E. coli

The procedures given below for production of biologically active, refolded TPO

25 (met-1 1-153) can be applied in analogy for the recovery of other TPO variants including N and C terminal extended forms (see Example 23).

A Recovery of non-soluble TPO (Met 1 1-153)

E. coli cells expressing TPO (Met<sup>-1</sup> 1-153) encoded by the plasmid pMP210-1 are fermented as described above. Typically, about 100g of cells are resuspended in 1 L (10 volumes) of cell disruption buffer (10 mM Tris, 5 mM EDTA, pH 8) with a Polytron homogenizer and the cells centrifuged at 5000 x g for 30 minutes. The washed cell pellet is again resuspended in 1 L cell disruption buffer with the Polytron homogenizer and the cell suspension is passed through an LH Cell Disrupter (LH inceltech, Inc.) or through a Microfluidizer (Microfluidics International) according to the manufactures' instructions. The suspension is centrifuged at 5,000g for 30 min. and resuspended and centrifuged a second time to make a washed refractile body pellet. The washed pellet is used immediately or stored frozen at -70°C.

# B. Solubilization and purification of monomeric TPO (Met 1 1-153)

The pellet from above is resuspended in 5 volumes by weight of 20 mM Tris, pH 8, with 6-8 M guanidine and 25 mM DTT (dithiothreitol) and stirred for 1-3 hr., or overnight, at 4°C to effect solubilization of the TPO protein. High concentrations of urea (6-8M) are also useful but generally result in lower yields compared to guanidine. After solubilization, the solution is centrifuged at 30,000 x g for 30 min. to produce a clear supernatant containing denatured, monomeric TPO protein. The supernatant is then chromatographed on a Superdex 200 gel filtration column (Pharmacia, 2.6 x 60 cm) at a flow rate of 2 ml/min. and the protein eluted with 20 mM Na phosphate, pH 6.0, with 10 mM DTT Fractions containing monomeric, denatured TPO protein eluting between 160 and 200 ml are pooled. The TPO protein is further purified on a semi-preparative C4 reversed phase column (2 x 20 cm VYDAC). The sample is applied at 5 ml/min. to a column equilibrated in 0.1% TFA(trifluoroacetic acid) with 30% acetonitrile. The protein is eluted with a linear gradient of acetonitrile (30-60% in 60 min.). The purified reduced protein elutes at approximately 50% acetonitrile. This material is used for refolding to obtain biologically active TPO variant.

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C. Generation of biologically active TPO (Met 1 1-153)

Approximately 20 mg of monomeric, reduced and denatured TPO protein in 40 ml 0.1% TFA/50% acetonitrile is diluted into 360 ml of refolding buffer containing optimally the following reagents:

50 mM Tris 0.3 M NaCl 5 mM EDTA

25 2% CHAPS detergent

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25% glycerol

5 mM oxidized glutathione

1 mM reduced glutathione

pH adjusted to 8.3

After mixing, the refolding buffer is gently stirred at 4°C for 12-48 hr to effect maximal refolding yields of the correct disulfide-bonded form of TPO (see below). The solution is then acidified with TFA to a final concentration of 0.2%, filtered through a 0.45 or 0.22 micron filter, and 1/10 volume of acetonitrile added. This solution is then pumped directly onto a C4 reversed phase column and the purified, refolded TPO (Met<sup>-1</sup> 1-153) eluted with the same gradient program as above. Refolded, biologically active TPO is eluted at approximately 45% acetonitrile under these conditions. Improper disulfide-bonded versions of TPO are eluted earlier.

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The final purified TPO (Met<sup>-1</sup> 1-153) is greater than 95% pure as assessed by SDS gels and analytical C4 reversed phase chromatography. For animal studies, the C4 purified material was dialyzed into physiologically compatible buffers. Isotonic buffers (10 mM Na acetate, pH 5.5, 10 mM Na succinate, pH 5.5 or 10 mM Na phosphate, pH 7.4) containing 150 mM NaCl and 0.01% Tween 80 were utilized.

Because of the high potency of TPO in the Ba/F3 assay (half maximal stimulation is achieved at approximately 3 pg/ml), it is possible to obtain biologically active material utilizing many different buffer, detergent and redox conditions. However, under most conditions only a small amount of properly folded material (<10%) is obtained. For commercial manufacturing processes, it is desirable to have refolding yields at least 10%, more preferably 30-50% and most preferably >50%. Many different detergents (Triton X-100, dodecyl-beta-maltoside, CHAPS, CHAPSO, SDS, sarkosyl, Tween 20 and Tween 80, Zwittergent 3-14 and others) were assessed for efficiency to support high refolding yields. Of these detergents, only the CHAPS family (CHAPS and CHAPSO) were found to be generally useful in the refolding reaction to limit protein aggregation and improper disulfide formation. Levels of CHAPS greater than 1% were most useful. Sodium chloride was required for best yields, with the optimal levels between 0.1 M and 0.5M. The presence of EDTA (1-5 mM) limited the amount of metal-catalyzed oxidation (and aggregation) which was observed with some preparations. Glycerol concentrations of greater than 15% produced the optimal refolding conditions. For maximum yields, it was essential to have both oxidized and reduced glutathione or oxidized and reduced cysteine as the redox reagent pair. Generally higher yields were observed when the mole ratio of oxidized reagent is equal to or in excess over the reduced reagent member of the redox pair. pH values between 7.5 and about 9 were optimal for refolding of these TPO variants. Organic solvents (e.g. ethanol, acetonitrile, methanol) were tolerated at concentrations of 10-15% or lower. Higher levels of organic solvents increased the amount of improperly folded forms. Tris and phosphate buffers were generally useful. Incubation at 4°C also produced higher levels of properly folded TPO.

Refolding yields of 40-60% (based on the amount of reduced and denatured TPO used in the refolding reaction) are typical for preparations of TPO that have been purified through the first C4 step. Active material can be obtained when less pure preparations (e.g. directly after the Superdex 200 column or after the initial refractile body extraction) although the yields are less due to extensive precipitation and interference of non-TPO proteins during the TPO refolding process.

Since TPO (Met<sup>-1</sup> 1-153) contains 4 cysteine residues, it is possible to generate three different disulfide versions of this protein:

version 1: disulfides between cysteine residues 1-4 and 2-3

version 2: disulfides between cysteine residues 1-2 and 3-4

version 3: disulfides between cysteine residues 1-3 and 2-4.

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During the initial exploration in determining refolding conditions, several different peaks containing the TPO protein were separated by C4 reversed phase chromatography. Only one of these peaks had significant biological activity as determined using the Ba/F3 assay. Subsequently, the refolding conditions were optimized to yield preferentially that version. Under these conditions, the misfolded versions are less than 10-20% of the total monomer TPO obtained.

The disulfide pattern for the biologically active TPO has been determined to be 1-4 and 2-3 by mass spectrometry and protein sequencing(i.e. version 1). Aliquots of the various C4-resolved peaks (5-10 nmoles) were digested with trypsin (1:25 mole ratio of trypsin to protein). The digestion mixture was analyzed by matrix-assisted laser desorption mass spectrometry before and after reduction with DTT. After reduction, masses corresponding to most of the larger tryptic peptides of TPO were detected. In the un-reduced samples, some of these masses were missing and new masses were observed. The mass of the new peaks corresponded basically to the sum of the individual tryptic peptides involved in the disulfide pair. Thus it was possible to unequivocally assign the disulfide pattern of the refolded, recombinant, biologically active TPO to be 1-4 and 2-3. This is consistent with the known disulfide pattern of the related molecule erythropoletin.

D. Biological activity of recombinant, refolded TPO (met 1-153)

Refolded and purified TPO (Met<sup>-1</sup> 1-153) has activity in both *in vitro* and *in vivo* assays. In the Ba/F3 assay, half-maximal stimulation of thymidine incorporation into the Ba/F3 cells was achieved at 3.3 pg /ml (0.3 pM). In the *mpl* receptor-based ELISA, half-maximal activity occurred at 1.9 ng/ml (120 pM). In normal and myelosuppressed animals produced by near-lethal X-radiation, TPO (Met<sup>-1</sup> 1-153) was highly potent (activity was seen at doses as low as 30 ng/mouse) to stimulate the production of new platelets.

30 EXAMPLE 23

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Production of Other Biologically Active TPO Variants in E. coli

Three different TPO variants produced in E. coli, purified and refolded into biological active forms are provided below.

(1) MLF - 13 residues from the bacterial-derived signal sequence STII are fused to the N-terminal domain of TPO (residues 1-155). The resulting sequence is <a href="MKKNIAFLLNAYASPAPPAC-----CVRRA">MKKNIAFLLNAYASPAPPAC-----CVRRA</a> (SEQ ID NO: 85)

where the leader sequence is underlined and C·····C represents Cys<sup>7</sup> through Cys<sup>151</sup>. This variant was constructed to provide a tyrosine for radio-iodination of TPO for receptor and biological studies.

(2) H8MLF - 7 residues from the STII sequence, 8 histidine residues and the Factor Xa enzymatic cleavage sequence IEGR are fused to the N-terminal domain (residues 1-155) of TPO. The sequence is

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MKKNIAFHHHHHHHHEGRSPAPPAC······CVRRA (SEQ ID NO: 86) where the leader sequence is underlined and C······C represents Cys<sup>7</sup> through Cys<sup>151</sup>. This variant, when purified and refolded, can be treated with the enzyme Factor Xa which will cleave after the arginine residue of the sequence IEGR yielding a TPO variant of 155 residues in length with a natural serine N-terminal amino acid.

(3) T-H8MLF - is prepared as described above for variant (2), except a thrombin sensitive sequence IEPR is fused to the N-terminal domain of TPO. The resulting sequence is

MKKNIAFHHHHHHHIEPRSPAPPAC·····CVRRA (SEQ ID NO: 87) where the leader sequence is underlined and and C·····C represents Cys<sup>7</sup> through Cys<sup>151</sup>. This variant, after purification and refolding can be treated with the enzyme thrombin to generate a natural N-terminal variant of TPO of 155 residues in length.

A Recovery, solubilization and purification of monomeric, biologically active TPO variants (1), (2), and (3).

All of the variants were expressed in E. coli. The majority of the variants were found in refractile bodies, as observed in Example 22 for TPO (Met<sup>-1</sup> 1-153). Identical procedures for the recovery, solubilization and purification of monomeric TPO variants was achieved as described in Example 22. Identical refolding conditions to those used for TPO (Met<sup>-1</sup> 1-153) were used with overall yields of 30-50%. After refolding, the TPO variants were purified by C4 reversed phase chromatography in 0.1% TFA utilizing an acetonitrile gradient as described previously. All of the TPO variants (in their unproteolyzed forms) had biological activity as assessed by the Ba/F3 assay, with half-maximal activities of 2-5 pM.

B. Proteolytic processing of Variants (2) and (3) to generate authentic N-terminal TPO (1-155).

TPO variants (2) and (3) above were designed with an enzymatically-cleavable leader peptide before the normal N-terminal amino acid residue of TPO. After refolding and purification of variants (2) and (3) as described above, each was subjected to digestion with the appropriate enzyme. For each variant, the acetonitrile from the C4 reversed phase step was removed by blowing a gentle stream of nitrogen

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on the solution. Thereafter the two variants were treated with either Factor Xa or thrombin as described below.

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For TPO variant (2), 1 M Tris buffer, pH 8, was added to the acetonitrile-free solution to a final concentration of 50 mM and the pH was adjusted to 8 if necessary. NaCl and CaCl2 were added to 0.1 M and 2 mM, respectively. Factor Xa (New England Biolabs) was added to achieve about a 1:25 to 1:100 mole ratio of enzyme to variant. The sample was incubated at room temperature for 1-2 hr. to achieve maximal cleavage as assessed by a change in migration on SDS gels representing the loss of the leader sequence. Thereafter, the reaction mixture was purified by C4 reversed phase chromatography using the same gradient and conditions as described above for the purification of properly folded variants. Uncleaved variant B was separated from cleaved variant (2) by these conditions. The N-terminal amino acids were shown to be SPAPP, indicating that removal of the N-terminal leader sequence was successful. Factor Xa also generated variable amounts of an internal cleavage within the TPO domain; cleavage was observed after the arginine residue at position number 118 generating an additional N-terminal sequence of TTAHKDP-(SEQ ID NO: 88). On nonreducing SDS gels, a single band at approximately 17000 daltons was observed for the Factor Xa cleaved variant; on reducing gels two bands were seen of molecular weight of approximately 12000 and 5000 daltons, consistent with cleavage at arginine 118. This observation also confirmed that the two parts of the molecule were held together by a disulfide bond between the 1st and 4th cysteine residues, as deduced from the tryptic digestion experiments described above. In the Ba/F3 biological assay, the purified TPO (1-155) variant, after removal of the N-terminal leader sequence and with the internal cleavage, had a half-maximal activity of 0.2 to 0.3 picomolar. The intact variant with the leader sequence had a half-maximal activity of 2-4 picomolar.

For variant (3), the digestion buffer consisted of 50 mM Tris, pH 8, 2% CHAPS, 0.3 M NaCl, 5 mM EDTA and human or bovine thrombin (Calbiochem) at a 1:25 to 1:50 by weight of enzyme to TPO variant protein. Digestion was conducted at room temperature for 2-6 hours. The progress of the digestion was assessed by SDS gels as described above for the Factor Xa cleavage reaction. Generally, more than 90% cleavage of the leader sequence was achieved in this time. The resultant TPO was purified on C4 reversed phase columns as described above and was shown to have the desired N-terminal by amino acid sequencing. Only very minor (<5%) amounts of an internal cleavage at the same arginine-threonine bond as observed above with Factor Xa was obtained. The resultant TPO protein had high biological activity with half-maximal responses in the Ba/F3 assay at 0.2-0.4 picomolar protein. In the mp/receptor based ELISA, this protein had a half-maximal response at 2-4 ng/ml purified protein (120-240 picomolar) while the intact variant containing the leader sequence

was less potent in both assays by 5-10 fold. For animal studies, the HPLC-purified cleaved protein was dialyzed into physiological acceptable buffers, with 150 mM NaCl, 0.01% Tween 80 and 10 mM sodium succinate, pH 5.5, or 10 mM sodium acetate, pH 5.5, or 10 mM sodium phosphate, pH 7.4. By HPLC and SDS gels, the purified protein was stable for several weeks when stored at 4°C. In normal and myelosuppressed mice, this purified TPO with the authentic N-terminal sequence was highly active, stimulating the production of platelets at doses as low as 30 ng/mouse.

### **EXAMPLE 24**

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### Synthetic mpl Ligand

Although Human *mpl* ligand (hML) is usually made using recombinant methods, it can also be synthesized via enzymatic ligation of synthetic peptide fragments using methods described below. Synthetic production of hML allows the incorporation of unnatural amino acids or synthetic functionalities such as polyethylene glycol. Previously, a mutant of the serine protease subtilisin BPN, subtiligase (S221C/P225A) was engineered to efficiently ligate peptide esters in aqueous solution (Abrahmsen *et al.*, *Biochem.*, 30:4151-4159 [1991]). It has now been shown that synthetic peptides can be enzymatically ligated in a sequential manor to produce enzymatically active long peptides and proteins such as ribonuclease A (Jackson *et al.*, *Science*, [1994]). This technology, described in more detail below, has enabled us to chemically synthesize long proteins that previously could be made only with recombinant DNA technology.

A general strategy for hML<sub>153</sub> synthesis using subtiligase is shown (Scheme 1). Beginning with a fully deprotected peptide corresponding to the C-terminal fragment of the protein, an N-terminal protected, C-terminal activated ester peptide is added along with subtiligase. When the reaction is complete, the product is isolated by reverse phase HPLC and the protecting group is removed from the N-terminus. The next peptide fragment is ligated, deprotected and the process is repeated using successive peptides until full length protein is obtained. The process is similar to solid phase methodology in that an N-terminal protected C-terminal activated petide is ligated to the N-terminus of the preceding peptide and protein is synthesized in a C->N direction. However because each coupling results in addition of up to 50 residues and the products are isolated after each ligation, much longer highly pure proteins can be synthesized in reasonable yields.

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Scheme 1. Strategy for Synthesis of hML Using Subtiligase

H<sub>2</sub>N-Peptide<sub>2</sub>-CO-NH-Peptide<sub>1</sub>-CO<sub>2</sub>

H<sub>2</sub>N-Peptide<sub>3</sub>-CO-NH-Peptide<sub>2</sub>-CO-NH-Peptide<sub>1</sub>-CO<sub>2</sub>

Based on our knowledge of the sequence specificity of the subtiligase as well as the amino acid sequence of the biologically active "epo-domain" of hML, we divided hML153 into seven fragments 18-25 residues in length. Test ligation tetrapeptides were synthesized to determine suitable ligation junctions for the 18-25mer's. **Table 13** shows the results of these test ligations.

### TABLE 13

hML Test Ligations. Donor and nucleophile peptides were dissolved at 10 mM in 100 mM tricine (pH 7.8) at 22°C. Ligase was added to a final concentration of 10  $\mu$ M from a 1.6 mg/mL stock (~70  $\mu$ M) and the ligation allowed to proceed overnight. Yields are based on % ligation vs. hydrolysis of the donor peptides.

Site	Donor (glc-K-NH <sub>2</sub> )	Nucleophile-NH <sub>2</sub>	%Hydrolysis	%Ligation
1 (23/24)	HVLH	SRLS	92	08
	(SEQ ID NO: 89)	(SEQ ID NO: 90)		
(22/23)	SHVL	HSRL	48	52
	(SEQ ID NO: 91)	(SEQ ID NO: 92)		
2 (46/47)	AVDF	SLGE	22	78
	(SEQ ID NO: 93)	(SEQ ID NO: 94)		
3 (69/70)	AVTL	LLEG	53	47
	(SEQ ID NO: 95)	(SEQ ID NO: 96)		
4 (89/90)	LSSL	LGQL	95	05
	(SEQ ID NO: 97)	(SEQ ID NO: 98)		
(88/89)	C(acm)LSS	LLGQ	00	00
	(SEQ ID NO: 99)	(SEQ ID NO: 100)		
(90/91)	SSLL	GQLS	45	55
	(SEQ ID NO: 101)	(SEQ ID NO: 102)		
(88/89)	CLSS	LLGQ	90	10
	(SEQ ID NO: 103)	(SEQ ID NO: 100)		
5(107/108)	LQSL	LGTQ	99	01
	(SEQ ID NO: 104)	(SEQ ID NO: 105)		
(106/107)	ALQS	LLGT	70	30
	(SEQ ID NO: 106)	(SEQ ID NO: 107)		
6(128/129)	NAIF	LSFQ	60	40
	(SEQ ID NO: 108)	(SEQ ID NO: 109)		

Based on these experiments, the ligation peptides indicated in **Table 14** should be efficiently ligated by the subtiligase. A suitable protecting group for the N-terminus of each donor ester peptide was needed to prevent self-ligation. We chose an isonicotinyl (iNOC) protecting group (Veber et al., J. Org. Chem., **42**:3286-3289 [1977]) because it is water soluble, it can be incorporated at the last step of solid phase peptide synthesis and it is stable to anhydrous HF used to deprotect and cleave peptides from the solid phase resin. In addition, it can be removed from the peptide after each ligation under mild reducing conditions (Zn/CH3CO2H) to afford a free N-terminus for subsequent ligations. A glycolate-lysyl-amide (glc-K-NH2) ester was used for C-terminal activation based on previous experiments which showed this to be efficiently acylated by subtiligase (Abrahmsen et al., Blochem., **30**:41:51-4159 [1991]). The iNOC-protected, glc-K-amide activated peptides can be synthesized

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using standard solid phase methods as outlined (Scheme 2). The peptides are then segentially ligated until the full protein is produced and the final product refolded in vitro. Based on homology with EPO, disulfide pairs are believed to be formed between cysteine residues 7 and 151 and between 28 and 85. Oxidation of the disulfides may be accomplished by simply stirring the reduced material under an oxygen atmosphere for several hours. The refolded material can then be purified by HPLC and fractions containing active protein pooled and lyophilized. As an alternative, disulfides can be differentially protected to control sequential oxidation between specific disulfide pairs. Protection of cysteines 7 and 151 with acetamidomethyl (acm) groups would ensure oxidation of 28 and 85. The acm groups could then be removed and residues 7 and 151 oxidized. Conversely, residues 28 and 85 could be acm protected and oxidized in case sequential oxidation is required for correct folding. Optoinally, Cysteins 28 and 85 may be substituted with another natural or unnatural residue other than Cys to insure proper oxidation of cysteins 7 and 151.

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### TABLE 14.

Peptide Fragments Used For Total Synthesis of h-ML Using Subtiligase

### <u>Fragment</u>

20 Sequence

1 (SEQ ID NO: 110)

iNOC-HN-SPAPPACDLRVLSKLLRDSHVL-glc-K-NH2 (1-22)

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2 (SEQ ID NO: 111)

INOC-HN-HSRLSQCPEVHPLPTPVLLPAVDF-glc-K-NH2 (23-46)

30 3 (SEQ ID NO: 112)

INOC-HN-SLGEWKTQMEETKAQDILGAVTL-glc-K-NH2 (47-69)

4 (SEQ ID NO: 113)

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INOC-HN-LLEGVMAARGQLGPTCLSSLL-glc-K-NH2 (70-90)

5 (SEQ ID NO: 114)

iNOC-HN-GQLSGQVRLLLGALQS-glc-K-NH2 (90-106)

5 6 (SEQ ID NO: 115)

iNOC-HN-LLGTQLPPQGRTTAHKDPNAIF-glc-K-NH2 (107-128)

7 (SEQ ID NO: 116)

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H<sub>2</sub>N-LSFQHLLRGKVRFLMLVGGSTLCVR-CO<sub>2</sub> (129-153)

Peptide ligations are carried out at 25°C in 100mM tricine, pH 8 (freshly prepared and degassed by vacuum filtration through a 5 μM filter). Typically the C-terminal fragment is dissolved in buffer (2-5 mM peptide) and a 10x stock solution of subtiligase (1 mg/ml in 100mM tricine, pH 8) is added to bring the final enzyme concentration to ~ 5μM. A 3-5 molar excess of the glc-K-NH<sub>2</sub> activated donor peptide is then added as a solid, dissolved, and the mixture allowed to stand at 25°C. The ligations are monitored by analytical reverse phase C18 HPLC (CH<sub>3</sub>CN/H<sub>2</sub>O gradient with 0.1% TFA). The ligation products are purified by preparative HPLC and lyophilized. Isonicotinyl (iNOC) deprotection was performed by stirring HCl activated zinc dust with the protected peptide in acetic acid. The zinc dust is removed by filtration and the acetic acid evaporated under vacuum. The resulting peptide can be used directly in the next ligation and the process is repeated. Synthetic hML<sub>153</sub> can be ligated by procedures analogous to those described above to synthetic or recombinant hML<sub>154-332</sub> to produce synthetic or semisynthetic full length hML.

Synthetic hML has many advantages over recombinant. Unnatural side chains can be introduced in order to improve potency or specificity. Polymer functionalities such as polyethylene glycol can be incorporated to improve duration of action. For example, polyethylene glycol can be attached to lysine residues of the individual fragments (Table 14) before or after one or more ligation steps have been performed. Protease sensitive peptide bonds can be removed or altered to improve stability *in vivo*. In addition, heavy atom derivatives can be synthesized to aid in structure determination.

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a) Lysyl-paramethylbenzhydrylamine (MBHA) resin 1 (0.63 meq./gm., Advanced ChemTech) is stirred with bromoacetic acid (5 eq.) and diisopropyl carbodiimide (5 eq.) for 1 h. at 25°C in dimethylacetamide (DMA) to afford the bromoacetyl derivative 2. b) The resin is washed extensively with DMA and individual Boc-protected amino acids (3 eq., Bachem) are esterified by stirring with sodium bicarbonate (6 eq.) in dimethylformamide (DMF) for 24 h. at 50°C to afford the corresponding glycolatephenylalanyl-amide-resin 3. The amino acetylated resin 3 is washed with DMF (3x) and dichloromethane (CH2Cl2) (3x) and can be stored at room temperature for several months. The resin 3 can then be loaded into an automated peptide synthesizer (Applied Biosystems 430A) and the peptides elongated using standard solid phase procedures (5). c) The N-α-Boc group is removed with a solution of 45% trifluoroacetic acid in CH<sub>2</sub>Cl<sub>2</sub>. d) Subsequent Boc-protected amino acids (5 eq.) are preactivated using benzotriazol-1-yl-oxy-tris-(dimethylamino) phosphonium hexafluorophosphate (BOP, 4 eq.) and N-methylmorpholine (NMM, 10 eq.) in DMA and coupled for 1-2 h. e) The final N- $\alpha$ -Boc group is removed (TFA/CH $_2$ Cl $_2$ ) to afford 4 and the isonicotinyl (iNOC) protecting group is introduced as described previously (4) via stirring with of

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4-isonicotinyl-2-4-dinitrophenyl carbonate (3 eq.) and NMM (6 eq.) in DMA at 25°C for 24 h. f) Cleavage and deprotection of the peptide via treatment with anhydrous HF (5% anisole/ 5% ethylmethyl sufide) at 0°C for 1 h. affords the iNOC-protected, glycolate-lys-amide activated peptide 5 which is purified by reverse phase C18 HPLC (CH<sub>3</sub>CN/H<sub>2</sub>O gradient, 0.1% TFA). The identity of all substrates is confirmed by mass spectrometry.

SUPPLEMENTAL ENABLEMENT

The invention as claimed is enabled in accordance with the above specification and readily available references and starting materials. Nevertheless, Applicants have deposited with the American Type Culture Collection, Rockville, Md., USA (ATCC) the cell line listed below:

Escherichia coli, DH10B-pBSK-hmpl | 1.8, ATCC accession no. CRL 69575, deposited February 24, 1994.

Plasmid, pSVI5.ID.LL.MLORF, ATCC accession no. CRL 75958, deposited December 2, 1994; and

CHO DP-12 cells, ML 1/50 MCB (labeled #1594), ATCC accession no. CRL 11770, deposited December 6, 1994.

This deposit was made under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and the regulations thereunder (Budapest Treaty). This assures maintenance of a viable culture for 30 years from date of deposit. The organisms will be made available by ATCC under the terms of the Budapest Treat, and subject to an agreement between Applicants and ATCC which assures unrestricted availability upon issuance of the pertinent U.S. patent. Availability of the deposited strain is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws.

\* \* \* \* \*

While the invention has necessarily been described in conjunction with preferred embodiments and specific working examples, one of ordinary skill, after reading the foregoing specification, will be able to effect various changes, substitutions of equivalents, and alterations to the subject matter set forth herein, without departing from the spirit and scope thereof. Hence, the invention can be practiced in ways other than those specifically described herein. It is therefore intended that the protection granted by letters patent hereon be limited only by the appended claims and equivalents thereof.

All references cited herein are hereby expressly incorporated by reference.

## SEQUENCE LISTING

	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: Genentech, Inc. Eaton, Dan L. de Sauvage, Frederic J.	٠.
4.0	(ii) TITLE OF INVENTION: THROMBOPOIETIN	:
10	(iii) NUMBER OF SEQUENCES: 144	
15	<ul> <li>(iv) CORRESPONDENCE ADDRESS:</li> <li>(A) ADDRESSEE: Genentech, Inc.</li> <li>(B) STREET: 460 Point San Bruno Blvd</li> <li>(C) CITY: South San Francisco</li> <li>(D) STATE: California</li> <li>(E) COUNTRY: USA</li> <li>(F) ZIP: 94080</li> </ul>	
20 25	<ul> <li>(v) COMPUTER READABLE FORM:</li> <li>(A) MEDIUM TYPE: 5.25 inch, 360 Kb floppy disk</li> <li>(B) COMPUTER: IBM PC compatible</li> <li>(C) OPERATING SYSTEM: PC-DOS/MS-DOS</li> <li>(D) SOFTWARE: patin (Genentech)</li> </ul>	
30	(vi) CURRENT APPLICATION DATA:  (A) APPLICATION NUMBER:  (B) FILING DATE:  (C) CLASSIFICATION:	
35	<pre>(vii) PRIOR APPLICATION DATA:     (A) APPLICATION NUMBER: 08/176553     (B) FILING DATE: 03-JAN-1994</pre>	
33	(vii) PRIOR APPLICATION DATA:  (A) APPLICATION NUMBER: 08/348657  (B) FILING DATE: 02-DEC-1994	
40	(vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: 08/185607 (B) FILING DATE: 21-JAN-1994	
45	<pre>(vii) PRIOR APPLICATION DATA:     (A) APPLICATION NUMBER: 08/348658     (B) FILING DATE: 02-DEC-1994</pre>	
50	<pre>(vii) PRIOR APPLICATION DATA:     (A) APPLICATION NUMBER: 08/196689     (B) FILING DATE: 15-FEB-1994</pre>	
55	<pre>(vii) PRIOR APPLICATION DATA:     (A) APPLICATION NUMBER: 08/223263     (B) FILING DATE: 04-APR-1994</pre>	
55	<pre>(vii) PRIOR APPLICATION DATA:   (A) APPLICATION NUMBER: 08/249376   (B) FILING DATE: 25-MAY-1994</pre>	

5	(viii	( <i>I</i>	A) N2 B) RI	AME: EGIS:	AGENT Wint TRAT: ENCE	cer, [ON 1	Dary NUMBI	yl B. ER: :	32,63		5PCT				
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	(2) I	NFOF	TAMS	ION I	FOR S	SEQ :	ID NO	0:1:							
15	(i	( <i>I</i>	A) LI 3) TY	engti Pe:	CHARA H: 35 amir DGY:	33 ar	mino cid		is						
	(xi	) SI	EQUE	NCE I	DESCI	RIPT	ION:	SEQ	ID 1	NO:1	:				
20	Met 1	Glu	Leu	Thr	Glu 5	Leu	Leu	Leu	Val	Val 10	Met	Leu	Leu	Leu	Thr 15
25	Ala .	Arg	Leu	Thr	Leu 20	Ser	Ser	Pro	Ala	Pro 25	Pro	Ala	Cys	Asp	Leu 30
	Arg	Val	Leu	Ser	Lys 35	Leu	Leu	Arg	Asp	Ser 40	His	Val	Leu	His	Ser 45
30	Arg	Leu	Ser	Gln	Cys 50	Pro	Glu	Val	His	Pro 55	Leu	Pro	Thr	Pro	Val 60
35	Leu	Leu	Pro	Ala	Val 65	Asp	Phe	Ser	Leu	Gly 70	Glu	Trp	Lys	Thr	Gln 75
	Met	Glu	Glu	Thr	Lys 80	Ala	Gln	Asp	Ile	Leu 85	Gly	Ala	Val	Thr	Leu 90
40	Leu	Leu	Glu	Gly	Val 95	Met	Ala	Ala	Arg	Gly 100	Gln	Leu	Gly	Pro	Thr 105
	Cys	Leu	Ser	Ser	Leu 110	Leu	Gly	Gln	Leu	Ser 115	Gly	Gln	Val	Arg	Leu 120
45	Leu	Leu	Gly		Leu 125	Gln	Ser	Leu	Leu	Gly 130	Thr	Gln	Leu	Pro	Pro 135
50	Gln	Gly	Arg	Thr	Thr 140	Ala	His	Lys	Asp	Pro 145	Asn	Ala	Ile	Phe	Leu 150
	Ser	Phe	Gln	His	Leu 155	Leu	Arg	Gly	Lys	Val 160	Arg	Phe	Leu	Met	Leu 165
55	Val	Gly	Gly	Ser	Thr 170	Leu	Суз	Val	Arg	Arg 175	Ala	Pro	Pro	Thr	Thr 180
	Ala	Val	Pro	Ser	Arg 185	Thr	Ser	Leu	Val	Leu 190	Thr	Leu	Asn	Glu	Leu 195
60	Pro .	Asn	Arg	Thr	Ser 200	Gly	Leu	Leu	G1u	Thr 205	Asn	Phe	Thr	Ala	Ser 210

:

	Ala	Arg	Thr	Thr	Gly 215	Ser	Gly	Leu	Leu	Lys 220	Trp	Gln	Gln	Gly	Phe 225		
5	Arg	Ala	Lys	Ile	Pro 230	Gly	Leu	Leu	Asn	Gln 235	Thr	Ser	Arg	Ser	Leu 240		
	Asp	Gln	Ile	Pro	Gly 245	Tyr	Leu	Asn	Arg	Ile 250	His	Glu	Leu	Leu	Asn 255		
10	Gly	Thr	Arg	Gly	Leu 260	Phe	Pro	Gly	Pro	Ser 265	Arg	Arg	Thr	Leu	Gly 270		
15	Ala	Pro	Asp	Ile	Ser 275	Ser	Gly	Thr	Ser	Asp 280	Thr	Gly	Ser	Leu	Pro 285		
	Pro	Asn	Leu	Gln	Pro 290		Tyr	Ser	Pro	Ser 295	Pro	Thr	His	Pro	Pro 300		
20	Thr	- Gly	Gln	Tyr	Thr 305		Phe	Pro	Lev	Pro 310	Pro	Thr	Leu	Pro	Thr 315		
	Pro	Val	. Val	Gln	Leu 320		Pro	Leu	Leu	325	Asr	Pro	Ser	Ala	Pro 330		
25	Thi	r Pro	Thr	Pro	Thr 335		Pro	Let	. Le	1 Asr 340	n Thi	: Sei	с Туг	Thr	His 345		
30	Se	r Gli	n Ası	ı Lev	Ser 350		ı Glu	1 Gly 35:	/ 3								
	(2)	INF	ORMA	rion	FOR	SEQ	ID 1	NO:2	:								
35	•		(A) : (B) ! (C) :	ENCE LENG TYPE STRAI TOPO	rh: : nu NDED	1795 clei NESS	bas c ac : si	es id	:								
40	(	xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	2:						
	TC	TTCC	TACC	CAT	CTGC	TCC	CĊAG	AGGG	CT G	CCTG	CTGT	G CA	.CTTG	GGTC	50		
45	Cī	GGAG	CCCT	TCT	CCAC	CCG	GATA	GATT.	CC I	CACC	CTTC	G CC	CGCC	TTTG	100		
	C	CCAC	CCTA	CTC	TGCC	CAG	AAGT	GCAA	GA G	CCTA	AGCC	G CC	CTCCA	TGGC	150		
50	C	CAGO	BAAGO	ATI	CAGG	GGA	GAGO	CCC	CAA A	ACAGO	GAG	C AC	:GCC#	AGCC	200		
55	G	ACAC(	CCGG	G CCA	\GAA7	GGA	GCT	BACT	BAA !	rtgct	CCT	CG TY	GTC!	ATGC7	250		
	T	CTCC'	raac:	r gcz	AAGG	TAA	CGC!	rgtc	CAG (	CCCG	GCTC	CT C	CTGC'	PTGT	300		
60	λ	<b>ርር</b> ሞር	CGAG	r cc	rcag'	raaa	CTG	CTTC	GTG .	ACTC	CCAT	GT C	CTTC	ACAG	350		

	AGACTGAGCC	AGTGCCCAGA	GGTTCACCCT	TTGCCTACAC	CTGTCCTGCT	400
5	GCCTGCTGTG	GACTTTAGCT	TGGGAGAATG	GAAAACCCAG	ATGGAGGAGA	450
	CCAAGGCACA	GGACATTCTG	GGAGCAGTGA	CCCTTCTGCT	GGAGGGAGTG	500
10	ATGGCAGCAC	GGGGACAACT	GGGACCCACT	TGCCTCTCAT	CCCTCCTGGG	550
15	GCAGCTTTCT	GGACAGGTCC	GTCTCCTCCT	TGGGGCCCTG	CAGAGCCTCC	600
	TTGGAACCCA	GCTTCCTCCA	CAGGGCAGGA	CCACAGCTCA	CAAGGATCCC	650
20	AATGCCATCT	TCCTGAGCTT	CCAACACCTG	CTCCGAGGAA	AGGTGCGTTT	700
	CCTGATGCTT	GTAGGAGGGT	CCACCCTCTG	CGTCAGGCGG	GCCCCACCCA	750
25	CCACAGCTGT	CCCCAGCAGA	ACCTCTCTAG	TCCTCACACT	GAACGAGCTC	800
30	CCAAACAGGA	CTTCTGGATT	GTTGGAGACA	AACTTCACTG	CCTCAGCCAG	850
	AACTACTGGC	TCTGGGCTTC	TGAAGTGGCA	GCAGGGATTC	AGAGCCAAGA	900
35	TTCCTGGTCT	GCTGAACCAA	ACCTCCAGGT	CCCTGGACCA	AATCCCCGGA	950
4.0	TACCTGAACA	GGATACACGA	ACTCTTGAAT	GGAACTCGTG	GACTCTTTCC	1000
40	TGGACCCTCA	CGCAGGACCC	TAGGAGCCCC	GGACATTTCC	TCAGGAACAT	1050
45	CAGACACAGG	CTCCCTGCCA	CCCAACCTCC	AGCCTGGATA	TTCTCCTTCC	1100
	CCAACCCATC	CTCCTACTGG	ACAGTATACG	CTCTTCCCTC	TTCCACCCAC	1150
50	CTTGCCCACC	CCTGTGGTCC	AGCTCCACCC	CCTGCTTCCT	GACCCTTCTG	1200
<b>.</b> .	CTCCAACGCC	CACCCCTACC	AGCCCTCTTC	TAAACACATC	CTACACCCAC	1250
55	TCCCAGAATC	TGTCTCAGGA	AGGGTAAGGT	TCTCAGACAC	TGCCGACATC	1300
60	AGCATTGTCT	CATGTACAGC	TCCCTTCCCT	GCAGGGCGCC	CCTGGGAGAC	1350

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	AACTGGACAA GATTTCCTAC TTTCTCCTGA AACCCAAAGC CCTGGTAAAA 1400
5	GGGATACACA GGACTGAAAA GGGAATCATT TTTCACTGTA CATTATAAAC 1450
Ū	CTTCAGAAGC TATTTTTTA AGCTATCAGC AATACTCATC AGAGCAGCTA 1500
10	GCTCTTTGGT CTATTTTCTG CAGAAATTTG CAACTCACTG ATTCTCTACA 1550
	TGCTCTTTTT CTGTGATAAC TCTGCAAAGG CCTGGGCTGG CCTGGCAGTT 1600
15	GAACAGAGGG AGAGACTAAC CTTGAGTCAG AAAACAGAGA AAGGGTAATT 1650
20	TCCTTTGCTT CAAATTCAAG GCCTTCCAAC GCCCCCATCC CCTTTACTAT 1700
	CATTCTCAGT GGGACTCTGA TCCCATATTC TTAACAGATC TTTACTCTTG 1750
25	AGAAATGAAT AAGCTTTCTC TCAGAAAAAA AAAAAAAAAA
30	(2) INFORMATION FOR SEQ ID NO:3:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 42 amino acids  (B) TYPE: amino acid
35	(D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
40	Leu Leu Leu Val Val Met Leu Leu Leu Thr Ala Arg Leu Thr Leu  1 5 10 15  Ser Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val Leu Ser Lys
	20 . 25
45	Leu Leu Arg Asp Ser His Val Leu His Ser Arg Leu 35 40 42
	(2) INFORMATION FOR SEQ ID NO:4:
50	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 390 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>
55	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
	GAATTCCTGG AATACCAGCT GACAATGATT TCCTCCTCAT CTTTCAACCT 50
60	CACCTCTCCT CATCTAAGAA TTGCTCCTCG TGGTCATGCT TCTCCTAACT 100

	GCAAGGCTAA CGCTGTCCAG CCCGGCTCCT CCTGCTTGTG ACCTCCGAGT 15
5	CCTCAGTAAA CTGCTTCGTG ACTCCCATGT CCTTCACAGC AGACTGGTGA 20
10	GAACTCCCAA CATTATCCCC TTTATCCGCG TAACTGGTAA GACACCCATA 25
	CTCCCAGGAA GACACCATCA CTTCCTCTAA CTCCTTGACC CAATGACTAT 30
15	TCTTCCCATA TTGTCCCCAC CTACTGATCA CACTCTCTGA CAAGAATTAT 35
20	TCTTCACAAT ACAGCCCGCA TTTAAAAGCT CTCGTCTAGA 390
20	
	(2) INFORMATION FOR SEQ ID NO:5:
25	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 390 bases</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
	CTTAAGGACC TTATGGTCGA CTGTTACTAA AGGAGGAGTA GAAAGTTGGA 50
35	GTGGAGAGGA GTAGATTCTT AACGAGGAGC ACCAGTACGA AGAGGATTGA 10
40	CGTTCCGATT GCGACAGGTC GGGCCGAGGA GGACGAACAC TGGAGGCTCA 15
	GGAGTCATTT GACGAAGCAC TGAGGGTACA GGAAGTGTCG TCTGACCACT 20
45	CTTGAGGGTT GTAATAGGGG AAATAGGCGC ATTGACCATT CTGTGGGTAT 25
	GAGGGTCCTT CTGTGGTAGT GAAGGAGATT GAGGAACTGG GTTACTGATA 30
50	AGAAGGGTAT AACAGGGGTG GATGACTAGT GTGAGAGACT GTTCTTAATA 35
55	AGAAGTGTTA TGTCGGGCGT AAATTTTCGA GAGCAGATCT 390
	(2) INFORMATION FOR SEQ ID NO:6:
60	(i) SEQUENCE CHARACTERISTICS:

		(I	3) T O) T	YPI	E: a OLOG	mino Y:	o ac line	id ar											
_								ON:											
5	Ser 1	Pro	Αla	a P	ro I	Pro 2	Ala	Cys	Asp	Le	u F	arg 10	Val	Le	eu :	Ser	Lys	Le 1	ս 5
10	Leu	Arg	Asp	, S	er E	His 20	Val	Leu	His	Se	r 1	Arg 25	Leu	Se	er (	Gln	Cys	Pr 3	0
	Glu	Val	Hi	s P	ro l	Leu 35	Pro	Thr	Pro	Va	1 1	Leu 40	Leu	P	ro	Ala	Val	As 4	.5
15	Phe	Ser	Le	u G	Sly (	Glu 50	Trp	Lys	Thr	· GI	ln I	Met 55	Glu	G	lu	Thr	Lys	Al 6	.a 50
	Gln	Asp	ıl.	e I	Ŀeu	Gly 65	Ala	Val	Thi	. Le	eu	Leu 70	Leu	G	lu	Gly	Val	Me	et 75
20	Ala	Ala	a Ar	g (	Sly	Gln 80	Leu	Gly	Pro	T]	hr	Cys 85	Lev	S	er	Ser	Leu	Le S	eu 90
25	Gly	Glr	ı Le	eu s	Ser	Gly 95	Gln	Val	Ar	g L	eu	Leu 100	Lev	1 6	3ly	Ala	Lev	1 G	ln 05
	Ser	Le	ı L∈	eu (	Gly	Thr 110	Gln	Leu	Pr	o P	ro	Gln 115	Gly	/ I	Arg	Thr	Thi	1	1a 20
30	His	Ly	s As	sp	Pro	Asn 125	Ala	Ile	. Ph	e L	eu	Ser 130	Pho	e (	Gln	His	Lei	1 1	eu 35
	Arg	G1;	у Гу	ys	Val	Arg 140		e Lev	ı Me	t L	eu	Val 145	G1	у (	Gly	Ser	Th	r L 1	eu 50
35	Cys	. Va	1 A:	rg	Arg	Ala 155		) Pro	Th	r I	hr	Ala 160	Va	1 :	Pro	Sei	r Ar	g 1 1	hr 65
40	Ser	. Le	eu V	al	Leu	Thr 170		ASI	n Gl	u I	Leu	Pro 175	As	n.	Arg	Thi	r Se	r (	81y 180
	Leu	ı Le	eu G	lu	Thr	Asr 185		e Th	r Al	la S	Ser	Ala 190	a Ar	g	Thr	Th	ŗ Gl	y 5	Ser 195
45	Gly	y L∈	eu L	eu	Lys	Trg 200		n Gl	n G	ly 1	Phe	209	Al 5	.a	Lys	s Il	e Pr	· · · ·	31y 210
	Le	u Le	eu A	sn <sub>.</sub>	Gln	Th:		r Ar	g S	er :	Lev	22	p G]	ın	Ile	e Pr	o Gl	У :	<b>Tyr</b> 225
50	Le	u As	sn A	ırg	Ιlε	Hi:		u Le	u L	eu .	Asr	Gl; 23	у Т1 5	nr	Arg	g Gl	y Le	eu :	Phe 240
55	Pr	o G	ly I	?ro	Ser	24		g Th	r L	eu	Gly	7 Al 25	a P: 0	ro	As	p Il	e Se	er	Ser 255
	G1	у Т	hr S	Ser	Ası	o Th 26		y Se	er L	eu	Pro	Pr 26	o A: 5	sn	Le	u Gl	n P	ro	Gly 270
60	• ту	r S	er 1	Pro	Se	r Pr 27		ır H	is P	ro	Pr	o Th 28	r G	lу	Gl	n Ty	/r T	hr	Leu 285

	Phe P	ro I	Leu	Pro	Pro 290	Thr	Leu	Pro	Thr	Pro 295	Val	Val	Gln	Leu	His 300
5	Pro L	eu I	Leu	Pro	Asp 305	Pro	Ser	Ala	Pro	Thr 310	Pro	Thr	Pro	Thr	Ser 315
10	Pro L	eu I	Leu	Asn	Thr 320	Ser	Tyr	Thr	His	Ser 325	Gln	Asn	Leu	Ser	Gln 330
	Glu G 3	32													
15	(2) IN														
	(1)	(A) (B)	LE TY	ICE C INGTH	: 16 amir	66 an	ino id		ls						
20	(xi)			POLO				SEQ	ID N	10:7:	:				
0.5	Ala P	ro I	Pro	Arg	Leu 5	Ile	Cys	Asp	Ser	Arg 10	Val	Leu	Glu	Arg	Tyr 15
25	Leu L	eu (	Glu	Ala	Lys 20	Glu	Ala	Glu	Asn	Ile 25	Thr	Thr	Gly	Cys	Ala 30
30	Glu H	lis (	Cys	Ser	Leu 35	Asn	Glu	Asn	Ile	Thr 40	Val	Pro	Asp	Thr	Lys 45
	Val A	sn I	Phe	Tyr	Ala 50	Trp	Lys	Arg	Met	Glu 55	Val	Gly	Gln	Gln	Ala 60
35	Val G	lu V	Val	Trp	Gln 65	Gly	Leu	Ala	Leu	Leu 70	Ser	Glu	Ala	Val	Leu 75
40	Arg G	ly (	3ln	Ala	Leu 80	Leu	Val	Asn	Ser	Ser 85	Gln	Pro	Trp	Glu	Pro 90
40	Leu G	ln I	Leu	His	Val 95	Asp	Lys	Ala	Val	Ser 100	Gly	Leu	Arg	Ser	Leu 105
45	Thr T	hr I	Leu							Gln 115		Glu	Ala	Ile	Ser 120
	Pro P	ro l	Asp	Ala	Ala 125	Ser	Ala	Ala	Pro	Leu 130	Arg	Thr	Ile	Thr	Ala 135
50	Asp T	hr I	Phe	Arg	Lys 140	Leu	Phe	Arg	Val	Tyr 145	Ser	Asn	Phe	Leu	Arg 150
55	Gly I	ys I	Leu	Lys	Leu 155	Tyr	Thr	Gly	Glu	Ala 160	Cys	Arg	Thr	Gly	Asp 165
	Arg 166														

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	(2) INFORMATION FOR SEQ ID NO:8:	
5	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 328 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
10	Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val Leu Ser Lys Leu 1 5 10 15	
	Leu Arg Asp Ser His Val Leu His Ser Arg Leu Ser Gln Cys Pro 20 25 30	
15	Glu Val His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala Val Asp 35 40 45	
20	Phe Ser Leu Gly Glu Trp Lys Thr Gln Met Glu Glu Thr Lys Ala 50 55 60	
	Gln Asp Ile Leu Gly Ala Val Thr Leu Leu Glu Gly Val Met 65 70 75	
25	Ala Ala Arg Gly Gln Leu Gly Pro Thr Cys Leu Ser Ser Leu Leu 80 85 90	
	Gly Gln Leu Ser Gly Gln Val Arg Leu Leu Leu Gly Ala Leu Gln 95 100 105	
30	Ser Leu Leu Gly Thr Gln Gly Arg Thr Thr Ala His Lys Asp Pro 110 115 120	
35	Asn Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly Lys Val 125 130 135	
	Arg Phe Leu Met Leu Val Gly Gly Ser Thr Leu Cys Val Arg Arg 140 145 150	
40	Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser Leu Val Leu 155 160 165	
	Thr Leu Asn Glu Leu Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr 170 175 180	
45	Asn Phe Thr Ala Ser Ala Arg Thr Thr Gly Ser Gly Leu Leu Lys 185 190 195	
50	Trp Gln Gln Gly Phe Arg Ala Lys Ile Pro Gly Leu Leu Asn Gln 200 205 210	
	Thr Ser Arg Ser Leu Asp Gln Ile Pro Gly Tyr Leu Asn Arg Ile 215 220 225	
55	His Glu Leu Leu Asn Gly Thr Arg Gly Leu Phe Pro Gly Pro Ser 230 235 240	
	Arg Arg Thr Leu Gly Ala Pro Asp Ile Ser Ser Gly Thr Ser Asp 245 250 255	
60	•	

	Thr	Gly	Ser	Leu	Pro 260	Pro	Asn	Leu	Gln	Pro 265	Gly	Tyr	Ser	Pro	Ser 270
5	Pro	Thr	His	Pro	Pro 275	Thr	Gly	Gln	Tyr	Thr 280	Leu	Phe	Pro	Leu	Pro 285
	Pro	Thr	Leu	Pro	Thr 290	Pro	Val	Val	Gln	Leu 295	His	Pro	Leu	Leu	Pro 300
10	Asp	Pro	Ser	Ala	Pro 305	Thr	Pro	Thr	Pro	Thr 310	Ser	Pro	Leu	Leu	Asn 315
15	Thr	Ser	Tyr	Thr	His 320	Ser	Gln	Asn	Leu	Ser 325	Gln	Glu	Gly 328		
	(2)	INFOF	TAMS	ON I	FOR S	SEQ I	D NO	0:9:							
20	(:	(E	4) LE 3) TY	NCE C ENGTH (PE: OPOLC	H: 26	55 ar	mino cid		ds			•			
	( <b>x</b> :	i) SE	EQUE	NCE I	DESCI	RIPT:	ON:	SEQ	ID 1	10:9	•				
25	Ser 1	Pro	Ala	Pro	Pro 5	Ala	Cys	Asp	Leu	Arg 10	Val	Leu	Ser	Lys	Leu 15
30	Leu	Arg	Asp	Ser	His 20	Val	Leu	His	Ser	Arg 25	Leu	Ser	Gln	Cys	Pro 30
50	Glu	Val	His	Pro	Leu 35	Pro	Thr	Pro	Val	Leu 40	Leu	Pro	Ala	Val	Asp 45
35	Phe	Ser	Leu	Gly	Glu 50	Trp	Lys	Thr	Gln	Met 55	Glu	Glu	Thr	Lys	Ala 60
	Gln	Asp	Ile	Leu	Gly 65	Ala	Val	Thr	Leu	Leu 70	Leu	Glu	Gly	Val	Met 75
40	Ala	Ala	Arg	Gly	Gln 80	Leu	Gly	Pro	Thr	Cys 85	Leu	Ser	Ser	Leu	Leu 90
45	Gly	Gln	Leu	Ser	Gly 95	Gln	Val	Arg	Leu	Leu 100	Leu	Gly	Ala	Leu	Gln 105
75	Ser	Leu	Leu	Gly	Thr 110	Gln	Leu	Pro	Pro	Gln 115	Gly	Arg	Tḥr	Thr	Ala 120
50	His	Lys	Asp	Pro	Asn 125	Ala	Ile	Phe	Leu	Ser 130	Phe	Gln	His	Leu	Leu 135
	Arg	Gly	Lys	Asp	Phe 140	Trp	Ile	Val	Gly	Asp 145	Lys	Leu -	His	Суз	Leu 150
55	Ser	Gln	Asn	Tyr	Trp 155	Leu	Trp	Ala	Ser	Glu 160		Ala	Ala	Gly	11e 165
60	Gln	Ser	Gln	Asp	Ser 170	Trp	Ser	Ala	Glu	Pro 175		Leu	Gln	Val	Pro 180

	Gly	Pro	Ası	n P		Arg :	Ile	Pro	Glu	Gln	Asp 190	Thr	Arg	Th	r L	eu (	Glu 195
5	Trp	Asn	Se	r T		Thr 200	Leu	Ser	Trp	Thr	Leu 205	Thr	Gln	As	p F	Pro	Arg 210
	Ser	Pro	G1	уН		Phe 215	Leu	Arg	Asn	Ile	Arg 220	His	Arg	Le	u E	Pro	Ala 225
10	Thr	Gln	Pr	o E		Ala 230	Trp	Ile	Phe	Ser	Phe 235	Pro	Asr	Pr	0 5	Ser	Ser 2 <b>4</b> 0
15	Tyr	Trp	Th	ır V	/al	Tyr 245	Ala	Leu	Pro	Ser	Ser 250	Thr	His	. Le	u 1	Ala	His 255
ıJ	Pro	Cys	: G1	Ly I		Ala 260	Pro	Pro	Pro	Ala	Ser 265						
20								ID NO		:							
	,		(A) (B)	LE TY	NGTH PE:	: 20 amii GY:	51 a no a	mino cid	aci	ds							
25											NO:1						
30	1					5					i Arg 10	)					15
			_			20					r Arg	5					30
35						35					40	)					Asp 45
						50			•		5!	5					Ala 60
40						65	•				7	0					Met 75
45						80	)				8	5					Leu 90
75						95	5				10	0					Gln 105
50	Se	r Le	eu I	Leu	Gly	Th:		n Gl	y Ar	g Th	r Th 11	r Al 5	.a H:	is I	ys	Ası	120
	As	n A	la:	Ile	Phe	Le: 12		r Ph	e Gl	n Hi	s Le 13		eu A	rg (	31y	Ly	3 Asp 135
55	Ph	e T	rp :	Ile	Val	14		p Ly	s Le	u Hi	.s Cy 14	rs Le	eu S	er (	31n	Ası	n Tyr 150
60	Tr	p L	eu '	Trp	Ala	a Se 15		u Va	1 A]	la Al	la G] 16	y I: 50	le G	ln :	Ser	: G1:	n Asp 165
60											. '						•

	Ser Trp Ser A	la Glu Pro 170	Asn Leu	Gln Val Pro 175	Gly Pro Asn Pr 18
5	Arg Ile Pro G	lu Gln Asr 185	Thr Arg	Thr Leu Glu 190	Trp Asn Ser Tr
	Thr Leu Ser T	rp Thr Leu 200	Thr Gln	Asp Pro Arg 205	Ser Pro Gly Hi 21
10	Phe Leu Arg A	sn Ile Arg 215	g His Arg	Leu Pro Ala 220	Thr Gln Pro Pr 22
15	Ala Trp Ile P	he Ser Phe 230	Pro Asn	Pro Ser Ser 235	Tyr Trp Thr Va 24
. •	Tyr Ala Leu P	ro Ser Ser 245	Thr His	Leu Ala His 250	Pro Cys Gly Pr 25
20	Ala Pro Pro P	ro Ala Ser 260 261			
	(2) INFORMATIO	N FOR SEQ	ID NO:11:		
25	(B) TYP (C) STR	E CHARACTE GTH: 7849 E: nucleic ANDEDNESS: OLOGY: lir	bases cacid single		
30	(xi) SEQUENC	E DESCRIPT	rion: seq	ID NO:11:	
	CCCAGCCTCC TT	TCTCTTGT T	rccctggtca	TGCCTGCCTC	CCTGTCTCCT 50
35	GTCTCTCCCT CC	CACACACA (	CCCACTATCC	TCCCAGCTAT	CCCTACACCC 100
40	TCCTTCCTAA TC	TTGGGAGA (	CATCTCGTCT	GGCTGGACGG	GAAAATTCCA 150
	GGATCTAGGC CA	CACTTCTC A	AGCAGACATG	CCCATCCTTG	GGGAGGAGGA 200
45	ACAGGAGAGA GC	CTGAGGAA (	GTTCTGGGGG	ACAGGGGGAT	GATGGGATCA 250
	AGGTCAGGCC AG	GAAGCCCC 1	rgaggacaga	GACTGTGGGG	AGACTGGGAC 300
50	TGGGAAGAAA GC	AAAGGAGC 1	ragagccagg	GCCAAAGGAA	AAGGGGGCC 350
5 5	AGCAGGGAGG TA	TTTGCGGG (	GGAGGTCCAG	CAGCTGTCTT	TCCTAAGACA 400
	GGGACACATG GG	CCTGGTTA 1	TTCCTCTTGT	CACATGTGGA	ACGGTAGGAG 450
60	ATGGAAGACG GA	GACAGAAC A	AGCAAAGGA	GGGCCCTTGGG	CACAGAGGTC 500

	TGTGTGTGTA	GCCATCCAAG	CCACTGGACC	CCAGCAGACG	AGCACCTAAG	220
5	CTCAGGCTTA	ACCCAGTGCA	CGTGTGCGCA	CATACATGTG	CCCCGCACCT	600
	GACAGTCCAC	TCAACCCGTC	CAAACCCTTT	CCCCATAACA	CCAACCCATA	650
10	ACAGGAGATT	TCTCTCATGT	GGGCAATATC	CGTGTTCCCA	CTTCGAAAGG	700
15	GGGAATGACA	AGATAGGACT	CCCTAGGGGA	TTACAGAAAG	AAAAGCAGGA	750
. •	AAGCAAGCAT	CCTGTTGGAT	TTCAGCAGCA	GGTATGATGT	CCAGGGAAAA	800
20	GAAATTTGGA	TAGCCAGGGA	GTGAAAACCC	CACCAATCTT	AAACAAGACC	850
	TCTGTGCTTC	TTCCCCAGCA	ACACAAATGT	CCTGCCAGAT	TCCTCCTGGA	900
25	AAAAAACTTC	TGCTCCTGTC	CCCCTCCAGG	TCCCAGGTTG	CCCATGTCCA	950
30	GGAAAAGATG	GATCCCCCTA	TCCAAATCTT	CTCCGTGGTG	TGTGTGGGTG	1000
	GAGGAGTGGA	CCCTGGTCCA	GGCAGGGGCT	CCAGGGAAGA	A GAAGGCGTCA	1050
35	CTTCCGGGGG	CCTTCACCAG	; TGTCTGGTGC	G CTCCCTTCTC	TGATTGGGC#	1100
	GAAGTGGCCC	AGGCAGGCG	T ATGACCTGC	r gctgtggag	GGCTGTGCC	1150
40	CACCGCCACA	TGTCTTCCT/	A CCCATCTGC	r ccccagagg	G CTGCCTGCTY	3 1200
45	TGCACTTGGG	TCCTGGAGC	CTTCTCCAC	C CGGTGAGTG	G CCAGCAGGG	г 1250
	GTGGGGTTAT	GTGAGGGTA	G AÁAGGACAG	C AAAGAGAAA	T GGGCTCCCA	3 1300
50	CTGGGGGAGG	GGCAGGCAA	A CTGGAACCT	A CAGGCACTG	A CCTTTGTCG	A 1350
	GAAGAGTGT	A GCCTTCCCA	G AATGGGAGG	A GCAGGGCAG	A GCAGGGGTA	G 1400
55	GGGGTGGGG	r GCTGGTTTC	T GAGGGACTG	A TCACTTACT	T GGTGGAATA	C 1450
60	AGCACAGCC	C TGGCTGGCC	C TAAGGAAAG	GGACATGAG	CCAGGGAGA	A 1500

	AATAAGAGAG	GGAGCTGCAC	TTAGGGCTTA	GCAAACACAG	TAGTAAGATG	1550
5	GACACAGCCC	CAATCCCCAT	TCTTAGCTGG	TCATTCCTCG	TTAGCTTAAG	1600
	GTTCTGAATC	TGGTGCTGGG	GAAGCTGGGC	CAGGCAAGCC	AGGGCGCAAG	1650
10	GAGAGGGTAA	TGGGAGGAGG	GCCCACTCAT	GTTGACAGAC	CTACAGGAAA	1700
	TCCCAATATT	GAATCAGGTG	CAAGCCTCTT	TGCACAACTT	GTGAAAGGAG	1750
1 5	GAGGAAGCCA	TGTGGGGGGT	CCTGTGAAGG	AACCGGAAGG	GGTTCTGCCA	1800
20	AGGGGGCAGG	GAGGCAGGTG	TGAGCTATGA	GACAGATATG	TTAGTGGGCG	1850
	CCTAAGACAA	GGTAAGCCCC	TAAGGTGGGC	ATCACCCAGC	AGGTGCCCGT	1900
25	TCCTGGGCAG	CTGGTCTCAG	GAAGGAAGTC	CCAGAACTGT	TAGCCCATCT	1950
	CTTGGCCTCA	GATAATGGAG	TATTTCAGGA	CTTGGAGTCC	AAAGAAAAGC	2000
30	TCCAGTGGCT	TTATGTGTGG	GGGTAGATAG	GGAAAGAATA	GAGGTTAATT	2050
35	TCTCCCATAC	CGCCTTTTAA	TCCTGACCTC	TAGTGGTCCC	AGTTACAGCT	2100
	TTGTGCAGTT	CCCCTCCCCA	GCCCACTCC	CCACCGCAGA	AGTTACCCCT	2150
40	CAACATATTG	CGCCCGTTTG	CCAGTTCCTC	ACCCAGGCCC	TGCATCCCAT	2200
4 5	TTTCCACTCT	CTTCTCCAGG	CTGAAGCCAC	AATACTTTCC	TTCTCTATCC	2250
45	CCATCCCAGA	TTTTCTCTGA	CCTAACAACC	AAGGTTGCTC	AGAATTŢAAG	2300
50	GCTAATTAAG	ATATGTGTGT	ATACATATCA	TGTCCTGCTG	CTCTCAGCAG	2350
	GGGTAGGTGG	CACCAAATCC	GTGTCCGATT	CACTGAGGAG	TCCTGACAAA	2400
5 5	AAGGAGACAC	CATATGCTTT	CTTGCTTTCT	TTCTTTCTTT	CTTTCTTTTT	2450
60	TTTTTTTGA	GACGGAGTTT	CACTCTTATT	GCCCAGGCTG	GAGTGCAATG	2500
	GTGCGATCTC	GGCTCACCAC	AAACCTCCGC	CTCCCAGGTA	CAAGCGATTC	2550

	TCCTGTCTCA GCCTCCCAAG TAGCTTGGAT TACAGGCATG AGCCACCACA 2600
5	CCCTGCTAGT TTTTTTGTAT TTCGTAGAGC CGGGGTTTCA CCATGTTAGT 2650
10	GAGGCTGGTG GCGAACTCCT GACCTCAGGT GATCCACCCG CCTTGGACTC 2700
	CCAAAGTGCT GGGATTACAG GCATGAGCCA CTGCACCCGG CACACCATAT 2750
15	GCTTTCATCA CAAGAAAATG TGAGAGAATT CAGGGCTTTG GCAGTTCCAG 2800
	GCTGGTCAGC ATCTCAAGCC CTCCCCAGCA TCTGTTCACC CTGCCAGGCA 2850
20	GTCTCTTCCT AGAAACTTGG TTAAATGTTC ACTCTTCTTG CTACTTTCAG 2900
25	GATAGATTCC TCACCCTTGG CCCGCCTTTG CCCCACCCTA CTCTGCCCAG 2950
	AAGTGCAAGA GCCTAAGCCG CCTCCATGGC CCCAGGAAGG ATTCAGGGGA 3000
30	GAGGCCCCAA ACAGGGAGCC ACGCCAGCCA GACACCCCGG CCAGAATGGA 3050
	GCTGACTGGT GAGAACACAC CTGAGGGGCT AGGGCCATAT GGAAACATGA 3100
35	CAGAAGGGGA GAGAGAAAGG AGACACGCTG CAGGGGGCAG GAAGCTGGGG 3150
40	GAACCCATTC TCCCAAAAAT AAGGGGTCTG AGGGGTGGAT TCCCTGGGTT 3200
	TCAGGTCTGG GTCCTGAATG GGAATTCCTG GAATACCAGC TGACAATGAT 3250
45	TTCCTCCTCA TCTTTCAACC TCACCTCTCC TCATCTAAGA ATTGCTCCTC 3300
	GTGGTCATGC TTCTCCTAAC TGCAAGGCTA ACGCTGTCCA GCCCGGCTCC 3350
50	TCCTGCTTGT GACCTCCGAG TCCTCAGTAA ACTGCTTCGT GACTCCCATG 3400
55	TCCTTCACAG CAGACTGGTG AGAACTCCCA ACATTATCCC CTTTATCCGC 3450
	GTAACTGGTA AGACACCCAT ACTCCCAGGA AGACACCATC ACTTCCTCTA 3500
60	ACTICITICAC CCAATGACTA TTCTTCCCAT ATTGTCCCCA CCTACTGATC 3550

	ACACTCTCTG	ACAAGAATTA	TTCTTCACAA	TACAGCCCGC	ATTTAAAAGC	3600
5	TCTCGTCTAG	AGATAGTACT	CATGGAGGAC	TAGCCTGCTT	ATTAGGCTAC	3650
	CATAGCTCTC	TCTATTTCAG	CTCCCTTCTC	CCCCCACCAA	TCTTTTTCAA	3700
10	CAGAGCCAGT	GCCCAGAGGT	TCACCCTTTG	CCTACACCTG	TCCTGCTGCC	3750
15	TGCTGTGGAC	TTTAGCTTGG	GAGAATGGAA	AACCCAGATG	GTAAGAAAGC	3800
	CATCCCTAAC	CTTGGCTTCC	CTAAGTCCTG	TCTTCAGTTT	CCCACTGCTT	3850
20	CCCATGGATT	CTCCAACATT	CTTGAGCTTT	TTAAAAATAT	CTCACCTTCA	3900
	GCTTGGCCAC	CCTAACCCAA	TCTACATTCA	CCTATGATGA	TAGCCTGTGG	3950
25	ATAAGATGAT	GGCTTGCAGG	TCCAATATGT	GAATAGATTT	GAAGCTGAAC	4000
30	ACCATGAAAA	GCTGGAGAGA	AATCGCTCAT	GGCCATGCCT	TTGACCTATT	4050
	CCYGTTCAGT	CTTCTTAAAT	TGGCATGAAG	AAGCAAGACT	CATATGTCAT	4100
35	CCACAGATGA	CACAAAGCTG	GGAAGTACCA	СТААААТААС	AAAAGACTGA	4150
40	ATCAAGATTC	AAATCACTGA	AAGACTAGGT	CAAAAACAAG	GTGAAACAAC	4200
40	AGAGATATAA	ACTTCTACAT	GTGGGCCGGG	GGCTCACGCC	TGTAATCCCA	4250
45	GCACTTTGGG	AGGCCGAGGC	AGGCAGATCA	CCTGAGGGCA	GGAGTTTGAG	4300
	AGCAGCCTGG	CCAACATGGC	GAAACCCCGT	CTCTACTAAG	AATACAAAAT	4350
50	TAGCCGGGCA	TGGTAGTGCA	TGCCTGTAAT	CCCAGCTACT	TGGAAGGCTG	4400
55	AAGCAGGAGA	ATCCCTTGAA	CCCAGGAGGT	GGAGGTTGTA	GTGAGCTGAG	4450
33	ATCATGCCAA	TGCACTCCAG	CCTGGGTGAC	AAGAGCAAAA	CTCCGTCTCA	4500
60	AAAAGAAAAA	AAAATTCTAC	ATGTGTAAAT	TAATGAGTAA	AGTCCTATTC	4550

	CAGCTTTCAG GCCACAATGC CCTGCTTCCA TCATTTAAGC CTCTGGCCCT 4600
5	AGCACTTCCT ACGAAAAGGA TCTGAGAGAA TTAAATTGCC CCCAAACTTA 4650
	CCATGTAACA TTACTGAAGC TGCTATTCTT AAAGCTAGTA ATTCTTGTCT 4700
10	GTTTGATGTT TAGCATCCCC ATTGTGGAAA TGCTCGTACA GAACTCTATT 4750
	CCGAGTGGAC TACACTTAAA TATACTGGCC TGAACACCGG ACATCCCCCT 4800
15	GAAGACATAT GCTAATTTAT TAAGAGGGAC CATATTAAAC TAACATGTGT 4850
20	CTAGAAAGCA GCAGCCTGAA CAGAAAGAGA CTAGAAGCAT GTTTTATGGG 4900
•	CAATAGTTTA AAAAACTAAA ATCTATCCTC AAGAACCCTA GCGTCCCTTC 4950
25	TTCCTTCAGG ACTGAGTCAG GGAAGAAGGG CAGTTCCTAT GGGTCCCTTC 5000
	TAGTCCTTTC TTTTCATCCT TATGATCATT ATGGTAGAGT CTCATACCTA 5050
30	CATTTAGTTT ATTTATTATT ATTATTTGAG ACGGAGTCTC ACTCTATCCC 5100
35	CCAGGCTGGA GTGCAGTGGC ATGATCTCAA CTCACTGCAA CCTCAGCCTC 5150
	CCGGATTCAA GCGATTCTCC TGCCTCAGTC TCCCAAGTAG CTGGGATTAC 5200
40	AGGTGCCCAC CACCATGCCC AGCTAATTTT TGTATTTTTG GTAGAGATGG 5250
	GGTTTCACCA TGTTGGCCAG GCTGATCTTG AACTCCTGAC CTCAGGTGAT 5300
45	CCACCTGCCT CAGCCTCCCA AAGTGCTGGG ATTACAGGCG TGAGCCACTG 5350
50	CACCCAGCCT TCATTCAGTT TAAAAATCAA ATGATCCTAA GGTTTTGCAG 5400
	CAGAAAGAGT AAATTTGCAG CACTAGAACC AAGAGGTAAA AGCTGTAACA 5450
55	GGGCAGATTT CAGCAACGTA AGAAAAAAGG AGCTCTTCTC ACTGAAACCA 5500
	AGTGTAAGAC CAGGCTGGAC TAGAGGACAC GGGAGTTTTT GAAGCAGAGG 5550
60	CTGATGACCA GCTGTCGGGA GACTGTGAAG GAATTCCTGC CCTGGGTGGG 5600

	ACCTTGGTCC	TGTCCAGTTC	TCAGCCTGTA	TGATTCACTC	TGCTGGCTAC	5650
5	TCCTAAGGCT	CCCCACCCGC	TTTTAGTGTG	CCCTTTGAGG	CAGTGCGCTT	5700
10	CTCTCTTCCA	TCTCTTTCTC	AGGAGGAGAC	CAAGGCACAG	GACATTCTGG	5750
	GAGCAGTGAC	CCTTCTGCTG	GAGGGAGTGA	TGGCAGCACG	GGGACAACTG	5800
15	GGACCCACTT	GCCTCTCATC	CCTCCTGGGG	CAGCTTTCTG	GACAGGTCCG	5850
•	TCTCCTCCTT	GGGGCCCTGC	AGAGCCTCCT	TGGAACCCAG	GTAAGTCCCC	5900
20	AGTCAAGGGA.	TCTGTAGAAA	CTGTTCTTTT	CTGACTCAGT	CCCACTAGAA	5950
25	GACCTGAGGG	AAGAAGGGCT	CTTCCAGGGA	GCTCAAGGGC	AGAAGAGCTG	6000
	ATCTACTAAG	AGTGCTCCCT	GCCAGCCACA	ATGCCTGGGT	ACTGGCATCC	6050
30	TGTCTTTCCT	ACTTAGACAA	GGGAGGCCTG	AGATCTGGCC	CTGGTGTTTG	6100
	GCCTCAGGAC	CATCCTCTGC	CCTCAGCTTC	CTCCACAGGG	CAGGACCACA	6150
35	GCTCACAAGG	ATCCCAATGC	CATCTTCCTG	AGTTTCCAAC	ACCTGCTCCG	6200
40	AGGAAAGGTG	CGTTTCCTGA	TGCTTGTAGG	AGGGTCCACC	CTCTGCGTCA	6250
	GCCGGCCCC	ACCCACCACA	GCTGTCCCCA	GCAGAACCTC	TCTAGTCCTC	6300
45	ACACTGAACG	AGCTCCCAAA	CAGGACTTCT	GGATTGTTGG	AGACAAACTT	6350
	CACTGCCTCA	GCCAGAACTA	CTGGCTCTGG	GCTTCTGAAG	TGGCAGCAGG	6400
50	GATTCAGAGC	CAAGATTCCT	GGTCTGCTGA	ACCAAACCTC	CAGGTCCCTG	6450
55	GACCAAATCC	CCGGATACCT	GAACAGGATA	CACGAACTCT	TGAATGGAAC	6500
	TCGTGGACTC	TTTCCTGGAC	CCTCACGCAG	GACCCTAGGA	GCCCCGGACA	6550
60	TTTCCTCAGG	AACATCAGAC	ACAGGCTCCC	TGCCACCCAA	CCTCCAGCCT	6600

:

	GGATATTCTC	CTTCCCCAAC	CCATCCTCCT	ACTGGACAGT	ATACGCTCTT	0030
5	CCCTCTTCCA	CCCACCTTGC	CCACCCCTGT	GGTCCAGCTC	CACCCCTGC	6700
	TTCCTGACCC	TTCTGCTCCA	ACGCCCACCC	CTACCAGCCC	TCTTCTAAAC	6750
10	ACATCCTACA	CCCACTCCCA	GAATCTGTCT	CAGGAAGGGT	AAGGTTCTCA	6800
15	GACACTGCCG	ACATCAGCAT	TGTCTCATGT	ACAGCTCCCT	TCCCTGCAGG	6850
. •	GCGCCCCTGG	GAGACAACTG	GACAAGATTT	CCTACTTTCT	CCTGAAACCC	6900
20	AAAGCCCTGG	TAAAAGGGAT	ACACAGGACT	GAAAAGGGAA	TCATTTTTCA	6950
	CTGTACATTA	TAAACCTTCA	GAAGCTATTT	TTTTAAGCTA	TCAGÇAATAC	7000
25	TCATCAGAGO	AGCTAGCTCT	TTGGTCTATT	TTCTGCAGAA	ATTTGCAACT	7050
30	CACTGATTCT	CTACATGCTC	TTTTTCTGTG	ATAACTCTGC	: AAAGGCCTGG	7100
00	GCTGGCCTGG	G CAGTTGAACA	A GAGGGAGAGA	A CTAACCTTG	GTCAGAAAAC	7150
35	AGAGAAAGG	TAATTTCCT	r TGCTTCAAA1	TCAAGGCCTT	CCAACGCCCC	7200
	CATCCCCTT	r actatcatt	TCAGTGGGA	TCTGATCCCA	A TATTCTTAAC	7250
40	AGATCTTTA	C TCTTGAGAA	A TGAATAAGC	r ttctctcag	A AATGCTGTCC	7300
45	CTATACACT	A GACAAAACT	G AGCCTGTATA	A AGGAATAAA'	r GGGAGCGCCC	3 7350
70	AAAAGCTCC	C TAAAAAGCA	A GGGAAAGAT	G TTCTTCGAG	G GTGGCAATA	3 7400
50	ATCCCCCTC	A CCCTGCCAC	C CCAAACAAA	A AAGCTAACA	G GAAGCCTTG	g <b>74</b> 50
	AGAGCCTCA	C ACCCCAGGT	A AGGCTGTGT	A GACAGTTCA	G TAAAGACAG	g 7500
55	ACCTGGATG	T GACAGCTGA	G CAAACAGCT	'A GAGCTTTGG	C AGCTCAGCA	G 7550
60	GAGGCTTTG	C CAGGCATGO	A CGCCTGCCT	C CCTCCTGTG	G AGGTCAGGA	G 7600

GAAGTGCAGG AAGTGGCATG AGTCAGGCTC CTTGAGCTCA CACAGCAGGA 7650 GAACAAGTAC AAGTCAAGTA CAAGTTGAAG GCTCATTTCC CAGTTCCCGC 7700 5 AAATGCATCT AAAAAGCAGC TCTGTGTGAC CACCATAAAC TCTGCTAGGG 7750 10 GATCTCTAAA AAGGAGTCAG GCTTATGGGG CTTTGCAAAT AAGTGCTGCC 7800 TTGGTGCTCA GGAAAAGGTT TGTGTTGCAC AAAACACAAA TTCCACTGC 7849 15 (2) INFORMATION FOR SEQ ID NO:12: (i) SEQUENCE CHARACTERISTICS: 20 (A) LENGTH: 1443 bases (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12: GAGTCCTTGG CCCACCTCTC TCCCACCCGA CTCTGCCGAA AGAAGCACAG 50 30 AAGCTCAAGC CGCCTCCATG GCCCCAGGAA AGATTCAGGG GAGAGGCCCC 100 ATACAGGGAG CCACTTCAGT TAGACACCCT GGCCAGAATG GAGCTGACTG 150 35 ATTTGCTCCT GGCGGCCATG CTTCTTGCAG TGGCAAGACT AACTCTGTCC 200 AGCCCCGTAG CTCCTGCCTG TGACCCCAGA CTCCTAAATA AACTGCTGCG 250 40 TGACTCCCAC CTCCTTCACA GCCGACTGAG TCAGTGTCCC GACGTCGACC 300 45 CTTTGTCTAT CCCTGTTCTG CTGCCTGCTG TGGACTTTAG CCTGGGAGAA 350 TGGAAAACCC AGACGGAACA GAGCAAGGCA CAGGACATTC TAGGGGCAGT 400 50 GTCCCTTCTA CTGGAGGGAG TGATGGCAGC ACGAGGACAG TTGGAACCCT 450 55 CCTGCCTCTC ATCCCTCCTG GGACAGCTTT CTGGGCAGGT TCGCCTCCTC 500 TTGGGGGCCC TGCAGGGCCT CCTAGGAACC CAGGGCAGGA CCACAGCTCA 550 60

CAAGGACCC AATGCCCTCT TCTTGAGCTT GCAACAACTG CTTCGGGGAA 600

	AGGTGCGCTT	CCTGCTTCTG	GTAGAAGGTC	CCACCCTCTG	TGTCAGACGG	650
5	ACCCTGCCAA	CCACAGCTGT	CCCAAGCAGT	ACTTCTCAAC	TCCTCACACT	700
10	AAACAAGTTC	CCAAACAGGA	CTTCTGGATT	GTTGGAGACG	AACTTCAGTG	750
	TCACAGCCAG	AACTGCTGGC	CCTGGACTTC	TGAGCAGGCT	TCAGGGATTC	800
15	AGAGTCAAGA	TTACTCCTGG	TCAGCTAAAT	CAAACCTCCA	GGTCCCCAGT	850
	CCAAATCTCT	GGATACCTGA	ACAGGACACA	CGGACCTGTG	AATGGAACTC	900
. 20	ATGGGCTCTT	TGCTGGAACC	TCACTTCAGA	CCCTGGAAGC	CTCAGACATC	950
25	TCGCCCGGAG	CTTTCAACAA	AGGCTCCCTG	GCATTCAACC	TCCAGGGTGG	1000
	ACTTCCTCCT	TCTCCAAGCC	TTGCTCCTG#	A TGGACACACA	CCCTTCCCTC	1050
30	CTTCACCTG	CTTGCCCACC	ACCCATGGAT	CTCCACCCCA	GCTCCACCC	1100
	CTGTTTCCT	ACCCTTCCA	CACCATGCC	r aactctacco	CCCCTCATCC	1150
35	AGTCACAATO	TACCCTCAT	C CCAGGAATT	T GTCTCAGGAA	A ACATAGCGCG	1200
40	GGCACTGGC	C CAGTGAGCG	T CTGCAGCTT	C TCTCGGGGA(	AAGCTTCCCC	1250
	AGGAAGGCT	G AGAGGCAGC	T GCATCTGCT	C CAGATGTTC	r GCTTTCACC	r 1300
45	AAAAGGCCC	T GGGGAAGGG	A TACACAGCA	C TGGAGATTG	ATTTTAAAA 1	3 1350
<b>5</b> 0	GAGCTATTT	т тттттаасс	T ATCAGCAAI	'A TTCATCAGA		A 1400
50	TCTTTGGTC	T ATTTTCGGT	A TAAATTTGA	A AATCACTAA	T TCT 1443	

- 55 (2) INFORMATION FOR SEQ ID NO:13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 352 amino acids
      (B) TYPE: amino acid
      (D) TOPOLOGY: linear
- 60

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

5	Met 1	Glu	Leu	Thr	Asp 5	Leu	Leu	Leu	Ala	Ala 10	Met	Leu	Leu	Ala	Val 15
	Ala	Arg	Leu	Thr	Leu 20	Ser	Ser	Pro	Val	Ala 25	Pro	Ala	Cys	Asp	Pro 30
10	Arg	Leu	Leu	Asn	Lys 35	Leu	Leu	Arg	Asp	Ser 40	His	Leu	Leu	His	Ser 45
	Arg	Leu	Ser	Gln	Cys 50	Pro	Asp	Val	Asp	Pro 55	Leu	Ser	Ile	Pro	Val 60
15	Leu	Leu	Pro	Ala	Val 65	Asp	Phe	Ser	Leu	Gly 70	Glu	Trp	Lys	Thr	Gln 75
20	Thr	Glu	Gln	Ser	80 80	Ala	Gln	Asp	Ile	Leu 85	Gly	Ala	Val	Ser	Leu 90
	Leu	Leu	Glu	Gly	Val 95	Met	Ala	Ala	Arg	Gly 100	Gln	Leu	Glu	Pro	Ser 105
25	Суѕ	Leu	Ser	Ser	Leu 110	Leu	Gly	Gln	Leu	Ser 115	Gly	Gln	Val	Arg	Leu 120
	Leu	Leu	Gly	Ala	Leu 125	Gln	Gly	Leu	Leu	Gly 130	Thr	Gln	Gly	Arg	Thr 135
30	Thr	Ala	His	Lys	Asp 140	Pro	Asn	Ala	Leu	Phe 145	Leu	Ser	Leu	Gln	Gln 150
35	Leu	Leu	Arg	Gly	Lys 155	Val	Arg	Phe	Leu	Leu 160	Leu	Val	Glu	Gly	Pro 165
	Thr	Leu	Cys	Val	Arg 170	Arg	Thr	Leu	Pro	Thr 175	Thr	Ala	Val	Pro	Ser 180
40	Ser	Thr	Ser	Gln	Leu 185	Leu	Thr	Leu	Asn	Lys 190	Phe	Pro	Asn	Arg	Thr 195
	Ser	Gly	Leu	Leu	Glu 200	Thr	Asn	Phe	Ser	Val 205	Thr	Ala	Arg	Thr	Ala 210
45	Gly	Pro	Gly	Leu	Leu 215	Ser	Arg	Leu	Gln	Gly 220		Arg	Val	Lys	Ile 225
50	Thr	Pro	Gly	Gln	Leu 230	Asn	Gln	Thr	Ser	Arg 235	Ser	Pro	Val	Gln	Ile 240
	Ser	Gly	Tyr	Leu	Asn 245	Arg	Thr	His	Gly	Pro 250	Val	Asn	Gly	Thr	His 255
55	Gly	Leu	Phe	Ala	Gly 260	Thr	Ser	Leu	Gln	Thr 265	Leu	Glu	Ala	Ser	Asp 270
	Ile	Ser	Pro	Gly	Ala 275	Phe	Asn	Lys	Gly	Ser 280	Leu	Ala	Phe	Asn	<b>Leu</b> 285
60	Gln	Gly	Gly	Leu	Pro 290	Pro	Ser	Pro	Ser	Leu 295	Ala	Pro	Asp	Gly	His 300

	Thr Pro Phe Pro Pro Ser Pro Ala Leu Pro Thr Thr His Gly Ser
5	Pro Pro Gln Leu His Pro Leu Phe Pro Asp Pro Ser Thr Thr Met 320 325 330
	Pro Asn Ser Thr Ala Pro His Pro Val Thr Met Tyr Pro His Pro 335 340 345
10	Arg Asn Leu Ser Gln Glu Thr 350 352
15	(2) INFORMATION FOR SEQ ID NO:14:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 1536 bases  (B) TYPE: nucleic acid
20	(C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
25	GAGTCCTTGG CCCACCTCTC TCCCACCCGA CTCTGCCGAA AGAAGCACAG 50
30	AAGCTCAAGC CGCCTCCATG GCCCCAGGAA AGATTCAGGG GAGAGGCCCC 100
	ATACAGGGAG CCACTTCAGT TAGACACCCT GGCCAGAATG GAGCTGACTG 150
35	ATTTGCTCCT GGCGGCCATG CTTCTTGCAG TGGCAAGACT AACTCTGTCC 200
	AGCCCCGTAG CTCCTGCCTG TGACCCCAGA CTCCTAAATA AACTGCTGCG 250
40	TGACTCCCAC CTCCTTCACA GCCGACTGAG TCAGTGTCCC GACGTCGACC 300
45	CTTTGTCTAT CCCTGTTCTG CTGCCTGCTG TGGACTTTAG CCTGGGAGAA 350
43	TGGAAAACCC AGACGGAACA GAGCAAGGCA CAGGACATTC TAGGGGCAGT 400
50	GTCCCTTCTA CTGGAGGGAG TGATGGCAGC ACGAGGACAG TTGGAACCCT 450
	CCTGCCTCTC ATCCCTCCTG GGACAGCTTT CTGGGCAGGT TCGCCTCCTC 500
55	TTGGGGGCCC TGCAGGGCCT CCTAGGAACC CAGCTTCCTC TACAGGGCAG 550
0.0	GACCACAGCT CACAAGGACC CCAATGCCCT CTTCTTGAGC TTGCAACAAC 600
60	TGCTTCGGGG AAAGGTGCGC TTCCTGCTTC TGGTAGAAGG TCCCACCCTC 650

	TGTGTCAGAC	GGACCCTGCC	AACCACAGCT	GTCCCAAGCA	GTACTTCTCA	700
5	ACTCCTCACA	CTAAACAAGT	TCCCAAACAG	GACTTCTGGA	TTGTTGGAGA	750
10	CGAACTTCAG	TGTCACAGCC	AGAACTGCTG	GCCCTGGACT	TCTGAGCAGG	800
	CTTCAGGGAT	TCAGAGTCAA	GATTACTCCT	GGTCAGCTAA	ATCAAACCTC	850
1 5	CAGGTCCCCA	GTCCAAATCT	CTGGATACCT	GAACAGGACA	CACGGACCTG	900
0.0	TGAATGGAAC	TCATGGGCTC	TTTGCTGGAA	CCTCACTTCA	GACCCTGGAA	950
20	GCCTCAGACA	TCTCGCCCGG	AGCTTTCAAC	AAAGGCTCCC	TGGCATTCAA	1000
25	CCTCCAGGGT	GGACTTCCTC	CTTCTCCAAG	CCTTGCTCCT	GATGGACACA	1050
	CACCCTTCCC	TCCTTCACCT	GCCTTGCCCA	CCACCCATGG	ATCTCCACCC	1100
30	CAGCTCCACC	CCCTGTTTCC	TGACCCTTCC	ACCACCATGC	CTAACTCTAC	1150
35	CGCCCTCAT	CCAGTCACAA	TGTACCCTCA	TCCCAGGAAT	TTGTCTCAGG	1200
	AAACATAGCG	CGGGCACTGG	CCCAGTGAGC	GTCTGCAGCT	TCTCTCGGGG	1250
40	ACAAGCTTCC	CCAGGAAGGC	TGAGAGGCAG	CTGCATCTGC	TCCAGATGTT	1300
	CTGCTTTCAC	CTAAAAGGCC	CTGGGGAAGG	GATACACAGC	ACTGGAGATT	1350
45	GTAAAATTTT	AGGAGCTATT	TTTTTTAAC	CTATCAGCAA	TATTCATCAG	1400
50	AGCAGCTAGC	GATCTTTGGT	CTATTTTCGG	TATAAATTTG	AAAATCACTA	1450
- •	AAAAAAAAA	АААААААА	ААААААААА	АААААААА	AAAAAAAA	1500
5 5	AAAAAAAA	AAAAAAAA	АААААААА	AAAAAA 1530	5	

- (2) INFORMATION FOR SEQ ID NO:15:
- 60 (i) SEQUENCE CHARACTERISTICS:
  (A) LENGTH: 356 amino acids

					amin GY:										
_	(xi	) SE	QUEN	ICE I	ESCR	IPTI	ON:	SEQ	ID N	0:15	:				
5	Met (	Glu	Leu	Thr	Asp 5	Leu	Leu	Leu	Ala	Ala 1	Met	Leu	Leu	Ala	Val 15
1 0	Ala	Arg	Leu	Thr	Leu 20	Ser	Ser	Pro	Val	Ala 25	Pro	Ala	Суз	Asp	Pro 30
	Arg	Leu	Leu	Asn	Lys 35	Leu	Leu	Arg	Asp	Ser 40	His	Leu	Leu	His	Ser 45
15	Arg	Leu	Ser	Gln	Cys 50	Pro	Asp	Val	Asp	Pro 55	Leu	Ser	Ile	Pro	Val 60
20					65					Gly 70					/5
					80					Leu 85				••	90
25					95					Gly 100					105
	_				110					Ser 115					120
30					125					Gly 130					135
35					140	)				145					Leu 150
					155	5				160					Leu 165
40					170	)				175					180
					185	5				190	)				195
45					200	0				205	)				Thr 210
50					215	5				220	)				225
					23	0				235	5				g Ser 240
55					24	5				250	)				o Val 255
					26	0				26	5				r Leu 270
60	Gl	u Al	a Se	er As	p I1 27		r Pr	o Gl	y Al	a Pho 28	e As O	n Ly	s Gl	y Se	r Leu 285

	Ala	Phe	Asn	Leu	Gln 290	Gly	Gly	Leu	Pro	Pro 295	Ser	Pro	Ser	Leu	Ala 300
5	Pro	Asp	Gly	His	Thr 305	Pro	Phe	Pro	Pro	Ser 310	Pro	Ala	Leu	Pro	Thr 315
10	Thr	His	Gly	Ser	Pro 320	Pro	Gln	Leu	His	Pro 325	Leu	Phe	Pro	Asp	Pro 330
	Ser	Thr	Thr	Met	Pro 335	Asn	Ser	Thr	Ala	Pro 340	His	Pro	Val	Thr	Met 345
15	Tyr	Pro	His	Pro	Arg 350	Asn	Leu	Ser	Gln	Glu 355					
	(2)	INFO	RMAT	ION I	FOR S	SEQ ]	ID NO	0:16:	:						
20	(:	(I	EQUEI A) LI B) T' O) TO	ENGTI (PE :	H: 24	ll ar no ac	mino cid		is						
	(x:	i) SI	EQUE	NCE I	DESCI	RIPT	ON:	SEQ	ID 1	NO:16	5:				
25	Ser 1	Pro	Val	Ala	Pro 5	Ala	Cys	Asp	Pro	Arg 10	Leu	Leu	Asn	Lys	Leu 15
30	Leu	Arg	Asp	Ser	His 20	Leu	Leu	His	Ser	Arg 25	Leu	Ser	Gln	Суз	Pro 30
	Asp	Val	Asp	Pro	Leu 35	Ser	Ile	Pro	Val	Leu 40	Leu	Pro	Ala	Val	Asp 45
35	Phe	Ser	Leu	Gly	<b>Glu</b> 50	Trp	Lys	Thr	Gln	Thr 55	Glu	Gln	Ser	Lys	Ala 60
40	Gln	Asp	Ile	Leu	Gly 65	Ala	Val	Ser	Leu	Leu 70	Leu	Glu	Gly	Val	Met 75
	Ala	Ala	Arg	Gly	Gln 80	Leu	Glu	Pro	Ser	Cys 85	Leu	Ser	Ser	Leu	Leu 90
45	Gly	Gln	Leu	Ser	Gly 95	Gln	Val	Arg	Leu	Leu 100	Leu	Gly	Ala	Leu	Gln 105
	Gly	Leu	Leu	Gly	Thr 110	Gln	Leu	Pro	Leu	Gln 115	Gly	Arg	Thr	Thr	Ala 120
50	His	Lys	Asp	Pro	Asn 125	Ala	Leu	Phe	Leu	Ser 130	Leu	Gln	Gln	Leu	Leu 135
55	Arg	Gly	Lys	Asp	Phe 140	Trp	Ile	Val	Gly	Asp 145	Glu	Leu	Gln	Cys	His 150
- •	Ser	Gln	Asn	Сув	Trp 155	Pro	Trp	Thr	Ser	Glu 160	Gln	Ala	Ser	Gly	Ile 165
60	Gln	Ser	Gln	Asp	Tyr 170	Ser	Trp	Ser	Ala	Lys 175	Ser	Asn	Leu	Gln	Val 180

	Pro S	Ser I	Pro A		Leu 185	Trp	Ile	Pro	Glu	Gln 190	Asp	Thr	Arg	Thr	Cys 195
5	Glu 7	rp i	Asn S	Ser '		Ala	Leu	Cys	Trp	Asn 205	Leu	Thr	Ser	Asp	Pro 210
	Gly S	Ser 1	Leu i		His 215	Leu	Ala	Arg	Ser	Phe 220	Gln	Gln	Arg	Leu	Pro 225
10	Gly :	Ile (	Gln :		Pro 230	Gly	Trp	Thr	Ser	Ser 235	Phe	Ser	Lys	Pro	Cys 240
15	Ser 241 (2) I	NFOR	MATI	on F	FOR S	SEQ I	D NO	): <b>17</b> :	:						
20	(i	(A (B	QUEN L) LE S) TY O) TO	NGTH PE:	I: 33 amir	35 an	nino cid	ICS: acio	is						
	(xi	) SE	QUEN	ICE I	DESCI	RIPT	ON:	SEQ	ID	NO:1	7:				
25	Ser 1	Pro	Val	Ala	Pro 5	Ala	Суз	Asp	Pro	Arg 10		Leu	ı Asr	Lys	Leu 15
	Leu	Arg	Asp	Ser	His 20	Leu	Leu	His	Ser	Arg 25		Ser	Glr	а Сув	Pro 30
30	Asp	Val	Asp	Pro	Leu 35	Ser	Ile	Pro	Val	Lev 40	Let	Pro	Ala	a Val	Asp 45
35	Phe	Ser	Leu	Gly	Glu 50		Lys	Thr	Glr	Thi 55	Glu	ı Glı	n Sei	r Lys	s Ala 60
	Gln	Asp	Ile	Leu	Gly 65		Val	Ser	Lev	1 Let 7(		ı Gl	u Gl	y Vai	Met 75
40	Ala	Ala	Arg	Gly	Gln 80		Glu	Pro	Se:	r Cys		ı Se	r Se	r Le	u Leu 90
	Gly	Gln	Leu	Ser	Gly 95		. Val	Arg	j Lei	u Lei 10	ı Le	u Gl	y Al	a Le	u Gln 105
45	Gly	Leu	Leu	Gly	Thr 110		. Le	ı Pro	Let	u Gl: 11		y Ar	g Th	r Th	r Ala 120
50	His	Lys	: Asp	Pro	Asr 125		ı Lei	ı Phe	e Le	u Se 13		u Gl	n Gl	n Le	u Leu 135
	Arg	Gly	, Lys	Val	140		e Lei	ı Le	ı Le	u Va 14		u Gl	y Pr	o Th	r Leu 150
55	Cys	: Val	Arg	Arg	Th:		ı Pr	o Th	r Th	r Al 16		1 Pr	o Se	er Se	r Thr 165
	Ser	Glr	n Leu	ı Leı	ı Th:		ı As:	n Ly	s Ph	e Pr 17		n Ar	g Tì	ır Se	er Gly 180
60															

	Leu	Leu	Glu	Thr	Asn 185	Phe	Ser	Val	Thr	Ala 190	Arg	Thr	Ala	Gly	Pro 195
5	Gly	Leu	Leu	Ser	Arg 200	Leu	Gln	Gly	Phe	Arg 205	Val	Lys	Ile	Thr	Pro 210
	Gly	Gln	Leu	Asn	Gln 215	Thr	Ser	Arg	Ser	Pro 220	Val	Gln	Ile	Ser	Gly 225
10	Tyr	Leu	Asn	Arg	Thr 230	His	Gly	Pro	Val	Asn 235	Gly	Thr	His	Gly	Leu 240
15	Phe	Ala	Gly	Thr	Ser 245	Leu	Gln	Thr	Leu	Glu 250	Ala	Ser	Asp	Ile	Ser 255
	Pro	Gly	Ala	Phe	Asn 260	Lys	Gly	Ser	Leu	Ala 265	Phe	Asn	Leu	Gln	Gly 270
20	Gly	Leu	Pro	Pro	Ser 275	Pro	Ser	Leu	Ala	Pro 280	Asp	Gly	His	Thr	Pro 285
	Phe	Pro	Pro	Ser	Pro 290	Ala	Leu	Pro	Thr	Thr 295	His	Gly	Ser	Pro	Pro 300
25	Gln	Leu	His	Pro	Leu 305	Phe	Pro	Asp	Pro	Ser 310	Thr	Thr	Met	Pro	Asn 315
30	Ser	Thr	Ala	Pro	His 320	Pro	Val	Thr	Met	Туr 325	Pro	His	Pro	Arg	Asn 330
	Leu	Ser	Gln	Glu	Thr 335										
35	(2) :	i) SI (1	EQUEI A) LI B) T	NCE ( ENGTI YPE:	CHARA H: 31 amin	ACTEI	RIST: mino cid	ICS:							
40	(x:	i) si	D) T( EQUEI					SEQ	ID 1	NO:18	3:				
45	Ser 1	Pro	Ala	Pro	Pro 5	Ala	Cys	Asp	Pro	Arg 10	Leu	Leu	Asn	Lys	Leu 15
	Leu	Arg	Asp	Ser	His 20	Val	Leu	His	Gly	Arg 25	Leu	Ser	Gln	Cys	Pro 30
50	Asp	Ile	Asn	Pro	Leu 35	Ser	Thr	Pro	Val	Leu 40	Leu	Pro	Ala	Val	Asp 45
	Phe	Thr	Leu	Gly	Glu 50	Trp	Lys	Thr	Gln	Thr 55	Glu	Gln	Thr	Lys	Ala 60
55	Gln	Asp	Val	Leu	Gly 65	Ala	Thr	Thr	Leu	Leu 70	Leu	Glu	Ala	Val	Met 75
60	Thr	Ala	Arg	Gly	Gln 80	Val	Gly	Pro	Pro	Суs 85	Leu	Ser	Ser	Leu	Leu 90

	Val	Gln	Leu	Ser	Gly 95	Gln	Val	Arg	Leu	Leu 100	Leu	Gly	Ala	Leu	Gln 105
5	Asp	Leu	Leu	Gly	Met 110	Gln	Leu	Pro	Pro	Gln 115	Gly	Arg	Thr	Thr	Ala 120
	His	Lys	Asp	Pro	Ser 125	Ala <sub>.</sub>	Ile	Phe	Leu	Asn 130	Phe	Gln	Gln	Leu	Leu 135
10	Arg	Gly	Lys	Val	Arg 140	Phe	Leu	Leu	Leu	Val 145	Val	Gly	Pro	Ser	Leu 150
15	Cys	Ala	Lys	Arg	Ala 155	Pro	Pro	Ala	Ile	Ala 160	Val	Pro	Ser	Ser	Thr 165
15	Ser	Pro	Phe	His	Thr 170	Leu	Asn	Lys	Leu	Pro 175	Asn	Arg	Thr	Ser	Gly 180
20	Leu	Leu	Glu	Thr	Asn 185	Ser	Ser	Ile	Ser	Ala 190	Arg	Thr	Thr	Gly	Ser 195
	Gly	Phe	Leu	Lys	Arg 200		Gln	Ala	Phe	Arg 205	Ala	Lys	Ile	Pro	Gly 210
25	Leu	ı Lev	. Asn	Gln	Thr 215		Arg	Ser	Lev	220	Gln	Ile	Pro	Gly	His 225
30	Glr	n Asr	Gly	Thr	His 230		Pro	Leu	Ser	Gly 235	Ile	His	: Gly	Leu	Phe 240
30	Pro	Gly	y Pro	Gln	245		Ala	Lev	Gly	/ Ala 250		) Asp	Ile	Pro	255
35	Ala	a Thi	r Sei	c Gly	260		ser Ser	Arg	Pro	265		r Lev	ı Glr	Pro	Gly 270
	Gl	u Se:	r Pro	o Ser	275		A His	Pro	Se:	280	o Gly	y Arg	у Туз	Thr	285
40	Ph	e Se	r Pr	o Sei	290		c Sei	e Pro	Se:	r Pro 29	o Th: 5	r Val	l Glr	ı Lev	300
45	Pr	o Le	u Le	u Pro	30		Se:	r Ala	a Il	e Th: 31	r Pr	o Ası	n Se	r Thi	r Ser 315
43	Pr	o Le	u Le	u Ph	e Ala 32		a Hi	s Pr	o Hi	s Ph 32	e G1 5	n As	n Le	u Se:	r Gln 330
50	G1	u G1 33													
	(2)	INF	'ORMA	TION	FOR	SEQ	ID :	NO:1	9:						
55		(i)	(B) (C)	ENCE LENG TYPE STRA TOPO	TH: : nu NDED	1026 clei NESS	bas c ac : si	es id ngle							
60		(xi)	SEQU	JENCE	DES	CRIF	TION	: SE	II Q	NO:	19:				

	AGCCCGGCTC	CICCIGCCIG	TGACCCCCGA	CICCIAAAIA	AACTGCTTCG	50
5	TGACTCCCAT	GTCCTTCACG	GCAGACTGAG	CCAGTGCCCA	GACATTAACC	100
	CTTTGTCCAC	ACCTGTCCTG	CTGCCTGCTG	TGGACTTCAC	CTTGGGAGAA	150
10	TGGAAAACCC	AGACGGAGCA	GACAAAGGCA	CAGGATGTCC	TGGGAGCCAC	200
15	AACCCTTCTG	CTGGAGGCAG	TGATGACAGC	ACGGGGACAA	GTGGGACCCC	250
	CTTGCCTCTC	ATCCCTGCTG	GTGCAGCTTT	CTGGACAGGT	TCGCCTCCTC	300
20	CTCGGGGCCC	TGCAGGACCT	CCTTGGAATG	CAGCTTCCTC	CACAGGGAAG	350
	GACCACAGCT	CACAAGGATC	CCAGTGCCAT	CTTCCTGAAC	TTCCAACAAC	400
25	TGCTCCGAGG	AAAGGTGCGT	TTCCTGCTCC	TTGTAGTGGG	GCCCTCCCTC	450
30	TGTGCCAAGA	GGGCCCCACC	CGCCATAGCT	GTCCCGAGCA	GCACCTCTCC	500
	ATTCCACACA	CTGAACAAGC	TCCCAAACAG	GACCTCTGGA	TTGTTGGAGA	550
35	CAAACTCCAG	TATCTCAGCC	AGAACTACTG	GCTCTGGATT	TCTCAAGAGG	600
40	CTGCAGGCAT	TCAGAGCCAA	GATTCCTGGT	CTGCTGAACC	AAACCTCCAG	650
40	GTCCCTAGAC	CAAATCCCTG	GACACCAGAA	TGGGACACAC	GGACCCTTGA	700
45	GTGGAATTCA	TGGACTCTTT	CCTGGACCCC	AACCCGGGGC	CCTCGGAGCT	750
	CCAGACATTC	CTCCAGCAAC	TTCAGGCATG	GGCTCCCGGC	CAACCTACCT	800
50	CCAGCCTGGA	GAGTCTCCTT	CCCCAGCTCA	CCCTTCTCCT	GGACGATACA	850
55	CTCTCTTCTC	TCCTTCACCC	ACCTCGCCCT	CCCCACAGT	CCAGCTCCAG	900
	CCTCTGCTTC	CTGACCCCTC	TGCGATCACA	CCCAACTCTA	CCAGTCCTCT	950
60	TCTATTTGCA	GCTCACCCTC	ATTTCCAGAA	CCTGTCTCAG	GAAGAGTAAG	1000

10

## GTGCTCAGAC CCTGCCAACT TCAGCA 1026

- 5 (2) INFORMATION FOR SEQ ID NO:20:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 1014 bases
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:
- 15 AGCCCGGCTC CTCCTGCCTG TGACCCCCGA CTCCTAAATA AACTGCTTCG 50 TGACTCCCAT GTCCTTCACG GCAGACTGAG CCAGTGCCCA GACATTAACC 100 20 CTTTGTCCAC ACCTGTCCTG CTGCCTGCTG TGGACTTCAC CTTGGGAGAA 150 TGGAAAACCC AGACGGAGCA GACAAAGGCA CAGGATGTCC TGGGAGCCAC 200 25 AACCCTTCTG CTGGAGGCAG TGATGACAGC ACGGGGACAA GTGGGACCCC 250 30 CTTGCCTCTC ATCCCTGCTG GTGCAGCTTT CTGGACAGGT TCGCCTCCTC 300 CTCGGGGCCC TGCAGGACCT CCTTGGAATG CAGGGAAGGA CCACAGCTCA 350 35 CAAGGATCCC AGTGCCATCT TCCTGAACTT CCAACAACTG CTCCGAGGAA 400 AGGTGCGTTT CCTGCTCCTT GTAGTGGGGC CCTCCCTCTG TGCCAAGAGG 450 40 GCCCCACCCG CCATAGCTGT CCCGAGCAGC ACCTCTCCAT TCCACACACT 500 45 GAACAAGCTC CCAAACAGGA CCTCTGGATT GTTGGAGACA AACTCCAGTA 550 TCTCAGCCAG AACTACTGGC TCTGGATTTC TCAAGAGGCT GCAGGCATTC 600 50 AGAGCCAAGA TTCCTGGTCT GCTGAACCAA ACCTCCAGGT CCCTAGACCA 650 AATCCCTGGA CACCAGAATG GGACACACGG ACCCTTGAGT GGAATTCATG 700 55 GACTCTTTCC TGGACCCCAA CCCGGGGCCC TCGGAGCTCC AGACATTCCT 750 60 CCAGCAACTT CAGGCATGGG CTCCCGGCCA ACCTACCTCC AGCCTGGAGA 800

	GTCTCCTTCC CCAGCTCACC CTTCTCCTGG ACGATACACT CTCTTCTCTC 850
. 5	CTTCACCCAC CTCGCCCTCC CCCACAGTCC AGCTCCAGCC TCTGCTTCCT 900
10	GACCCCTCTG CGATCACACC CAACTCTACC AGTCCTCTTC TATTTGCAGC 950
	TCACCCTCAT TTCCAGAACC TGTCTCAGGA AGAGTAAGGT GCTCAGACCC 1000
15	TGCCAACTTC AGCA 1014
20	(i) SEQUENCE CHARACTERISTICS:
25	(A) LENGTH: 328 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear
23	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21: Ser Pro Ala Pro Pro Ala Cys Asp Pro Arg Leu Leu Asn Lys Leu
30	1 5 10 15
	20 25 30  Asp Ile Asn Pro Leu Ser Thr Pro Val Leu Leu Pro Ala Val Asp
35	35 40 45  Phe Thr Leu Gly Glu Trp Lys Thr Gln Thr Glu Gln Thr Lys Ala
40	The same of the sa
	65 70 75  Thr Ala Arg Gly Gln Val Gly Pro Pro Cys Leu Ser Ser Leu Leu 80 85 90
45	
50	Asp Leu Leu Gly Met Gln Gly Arg Thr Thr Ala His Lys Asp Pro
	Ser Ala Ile Phe Leu Asn Phe Gln Gln Leu Leu Arg Gly Lys Val
55	Arg Phe Leu Leu Val Val Gly Pro Ser Leu Cys Ala Lys Arg 140 145 150
60	Ala Pro Pro Ala Ile Ala Val Pro Ser Ser Thr Ser Pro Phe His 155 160 165

Thr Leu Asn Lys Leu Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr 170 175 180	
Asn Ser Ser Ile Ser Ala Arg Thr Thr Gly Ser Gly Phe Leu Lys 185 190 195	•
Arg Leu Gln Ala Phe Arg Ala Lys Ile Pro Gly Leu Leu Asn Gln 200 205 210	•
Thr Ser Arg Ser Leu Asp Gln Ile Pro Gly His Gln Asn Gly Thr 215 220 225	<b>.</b>
His Gly Pro Leu Ser Gly Ile His Gly Leu Phe Pro Gly Pro Gln 230 235 240	
Pro Gly Ala Leu Gly Ala Pro Asp Ile Pro Pro Ala Thr Ser Gly 245 250 255	
Met Gly Ser Arg Pro Thr Tyr Leu Gln Pro Gly Glu Ser Pro Ser 260 265 270	
Pro Ala His Pro Ser Pro Gly Arg Tyr Thr Leu Phe Ser Pro Ser 275 280 285	
Pro Thr Ser Pro Ser Pro Thr Val Gln Leu Gln Pro Leu Leu Pro 290 295 300	
Asp Pro Ser Ala Ile Thr Pro Asn Ser Thr Ser Pro Leu Leu Phe 305 310 315	
Ala Ala His Pro His Phe Gln Asn Leu Ser Gln Glu Glu 320 325 328	
(2) INFORMATION FOR SEQ ID NO:22:	
(A) LENGTH: 5141 bases (B) TYPE: nucleic acid	
(D) TOPOLOGY: linear	
•	
	Asn Ser Ser Ile Ser Ala Arg Thr Thr Gly Ser Gly Phe Leu Lys 195  Arg Leu Gln Ala Phe Arg Ala Lys Ile Pro Gly Leu Leu Asn Gln 200 205 210  Thr Ser Arg Ser Leu Asp Gln Ile Pro Gly His Gln Asn Gly Thr 215 220 225  His Gly Pro Leu Ser Gly Ile His Gly Leu Phe Pro Gly Pro Gln 230 240  Pro Gly Ala Leu Gly Ala Pro Asp Ile Pro Pro Ala Thr Ser Gly 245 255  Met Gly Ser Arg Pro Thr Tyr Leu Gln Pro Gly Glu Ser Pro Ser 265 270  Pro Ala His Pro Ser Pro Gly Arg Tyr Thr Leu Phe Ser Pro Ser 275  Pro Thr Ser Pro Ser Pro Thr Val Gln Leu Gln Pro Leu Leu Pro 290 300  Asp Pro Ser Ala Ile Thr Pro Asn Ser Thr Ser Pro Leu Leu Phe 315  Ala Ala His Pro His Phe Gln Asn Leu Ser Gln Glu Glu 320 328  (2) INFORMATION FOR SEQ ID NO:22:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: S141 bases (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:  TTCGAGCTCG CCCGACATTG ATTATTGACT AGAGTCGATC GACAGCTGTO 50  GAAGTATGCA AAGCATGCAT CTCAATTAGT CAGCAACCAG GTGTGGAAAG 150  TCCCCCAGGCT CCCCCAGCAGG CAGAAGTATG CAAAGCATGC ATCTCAATTA 200  GTCAGCAACC ATAGTCCCGC CCCTAACTCC GCCCATCCCG CCCCTAACTC 250

	TATGCAGAGG	CCGAGGCCGC	CTCGGCCTCT	GAGCTATTCC	AGAAGTAGTG	350
5	AGGAGGCTTT	TTTGGAGGCC	TAGGCTTTTG	CAAAAAGCTA	GCTTATCCGG	400
	CCGGGAACGG	TGCATTGGAA	CGCGGATTCC	CCGTGCCAAG	AGTGACGTAA	450
10	GTACCGCCTA	TAGAGCGATA	AGAGGATTTT	ATCCCCGCTG	CCATCATGGT	500
15	TCGACCATTG	AACTGCATCG	TCGCCGTGTC	CCAAAATATG	GGGATTGGCA	550
	AGAACGGAGA	CCTACCCTGG	CCTCCGCTCA	GGAACGAGTT	CAAGTACTTC	600
20	CAAAGAATGA	·	TTCAGTGGAA	GGTAAACAGA	ATCTGGTGAT	650
	TATGGGTAGG	AAAACCTGGT	TCTCCATTCC	TGAGAAGAAT	CGACCTTTAA	700
25	AGGACAGAAT	TAATATAGTT	CTCAGTAGAG	AACTCAAAGA	ACCACCACGA	750
30	GGAGCTCATT	TTCTTGCCAA	AAGTTTGGAT	GATGCCTTAA	GACTTATTGA	800
	ACAACCGGAA	TTGGCAAGTA	AAGTAGACAT	GGTTTGGATA	GTCGGAGGCA	850
35	GTTCTGTTTA	CCAGGAAGCC	ATGAATCAAC	CAGGCCACCT	TAGACTCTTT	900
40	GTGACAAGGA	TCATGCAGGA	ATTTGAAAGT	GACACGTTTT	TCCCAGAAAT	950
70	TGATTTGGGG	AAATATAAAC	CTCTCCCAGA	ATACCCAGGC	GTCCTCTCTG	1000
45	AGGTCCAGGA	GGAAAAAGGC	ATCAAGTATA	AGTTTGAAGT	CTACGAGAAG	1050
	AAAGACTAAC	AGGAAGATGC	TTTCAAGTTC	TCTGCTCCCC	TCCTAAAGCT	1100
50	ATGCATTTTT	ATAAGACCAT	GGGACTTTTG	CTGGCTTTAG	ATCCCCTTGG	1150
55	CTTCGTTAGA	ACGCGGCTAC	AATTAATACA	TAACCTTATG	TATCATACAC	1200
	ATACGATTTA	GGTGACACTA	TAGATAACAT	CCACTTTGCC	TTTCTCTCCA	1250
60	CAGGTGTCCA	CTCCCAGGTC	CAACTGCACC	TCGGTTCTAA	GCTTCTGCAG	1300

	GTCGACTCTA GAGGATCCCC GGGGAATTCA ATCGATGGCC GCCATGGCCC 1350
5	AACTTGTTTA TTGCAGCTTA TAATGGTTAC AAATAAAGCA ATAGCATCAC 1400
	AAATTTCACA AATAAAGCAT TTTTTTCACT GCATTCTAGT TGTGGTTTGT 1450
10	CCAAACTCAT CAATGTATCT TATCATGTCT GGATCGATCG GGAATTAATT 1500
	CGGCGCAGCA CCATGGCCTG AAATAACCTC TGAAAGAGGA ACTTGGTTAG 1550
15	GTACCTTCTG AGGCGGAAAG AACCAGCTGT GGAATGTGTG TCAGTTAGGG 1600
20	TGTGGAAAGT CCCCAGGCTC CCCAGCAGGC AGAAGTATGC AAAGCATGCA 1650
	TCTCAATTAG TCAGCAACCA GGTGTGGAAA GTCCCCAGGC TCCCCAGCAG 1700
25	GCAGAAGTAT GCAAAGCATG CATCTCAATT AGTCAGCAAC CATAGTCCCG 1750
	CCCCTAACTC CGCCCATCCC GCCCCTAACT CCGCCCAGTT CCGCCCATTC 1800
30	TCCGCCCCAT GGCTGACTAA TTTTTTTTAT TTATGCAGAG GCCGAGGCCG 1850
35	CCTCGGCCTC TGAGCTATTC CAGAAGTAGT GAGGAGGCTT TTTTGGAGGC 1900
	CTAGGCTTTT GCAAAAAGCT GTTACCTCGA GCGGCCGCTT AATTAAGGCG 1950
40	CGCCATTTAA ATCCTGCAGG TAACAGCTTG GCACTGGCCG TCGTTTTACA 2000
	ACGTCGTGAC TGGGAAAACC CTGGCGTTAC CCAACTTAAT CGCCTTGCAG 2050
45	CACATCCCCC CTTCGCCAGC TGGCGTAATA GCGAAGAGGC CCGCACCGAT 2100
50	CGCCCTTCCC AACAGTTGCG TAGCCTGAAT GGCGAATGGC GCCTGATGCG 2150
	GTATTTTCTC CTTACGCATC TGTGCGGTAT TTCACACCGC ATACGTCAAA 2200
55	GCAACCATAG TACGCGCCCT GTAGCGGCGC ATTAAGCGCG GCGGGTGTGG 2250
	TGGTTACGCG CAGCGTGACC GCTACACTTG CCAGCGCCCT AGCGCCCCGCT 2300
60	CCTTTCGCTT TCTTCCCTTC CTTTCTCGCC ACGTTCGCCG GCTTTCCCCG 2350

		TCAAGCTCTA	AATCGGGGGC	TCCCTTTAGG	GTTCCGATTT	AGTGCTTTAC	2400
	5	GGCACCTCGA	CCCCAAAAAA	CTTGATTTGG	GTGATGGTTC	ACGTAGTGGG	2450
	10	CCATCGCCCT	GATAGACGGT	TTTTCGCCCT	TTGACGTTGG	AGTCCACGTT	2500
		CTTTAATAGT	GGACTCTTGT	TCCAAACTGG	AACAACACTC	AACCCTATCT	2550
•	15	CGGGCTATTC	TTTTGATTTA	TAAGGGATTT	TGCCGATTTC	GGCCTATTGG	2600
	•	TTAAAAAATG	AGCTGATTTA	ACAAAAATTT	AACGCGAATT	ТТААСААААТ	2650
	20	ATTAACGTTT	ACAATTTTAT	GGTGCACTCT	CAGTACAATC	TGCTCTGATG	2700
	25	CCGCATAGTT	AAGCCAACTC	CGCTATCGCT	ACGTGACTGG	GTCATGGCTG	2750
		CGCCCCGACA	CCCGCCAACA	CCCGCTGACG	CGCCCTGACG	GGCTTGTCTG	2800
	30	CTCCCGGCAT	CCGCTTACAG	ACAAGCTGTG	ACCGTCTCCG	GGAGCTGCAT	2850
	35	GTGTCAGAGG	TTTTCACCGT	CATCACCGAA	ACGCGCGAGG	CAGTATTCTT	2900
	33	GAAGACGAAA	GGGCCTCGTG	ATACGCCTAT	TTTTATAGGT	TAATGTCATG	2950
	40	ATAATAATGG	TTTCTTAGAC	GTCAGGTGGC	ACTTTTCGGG	GAAATGTGCG	3000
		CGGAACCCCT	ATTTGTTTAT	TTTTCTAAAT	ACATTCAAAT	ATGTATCCGC	3050
	45	TCATGAGACA	ATAACCCTGA	TAAATGCTTC	AATAATATTG	AAAAAGGAAG	3100
	50	AGTATGAGTA	TTCAACATTT	CCGTGTCGCC	CTTATTCCCT	TTTTTGCGGC	3150
_		ATTTTGCCTT	CCTGTTTTTG	CTCACCCAGA	AACGCTGGTG	AAAGTAAAAG	3200
	55	ATGCTGAAGA	TCAGTTGGGT	GCACGAGTGG	GTTACATCGA	ACTGGATCTC	3250
•		AACAGCGGTA	AGATCCTTGA	GAGTTTTCGC	CCCGAAGAAC	GTTTTCCAAT	33,00
	60	CAMCACCACM	mmm x x cmmc	mooma moneo	CCCCCMARMA	moocomos mo	2250

	ACGCCGGGCA	AGAGCAACTC	GGTCGCCGCA	TACACTATTC	TCAGAATGAC	3400
5	TTGGTTGAGT	ACTCACCAGT	CACAGAAAAG	CATCTTACGG	ATGGCATGAC	3450
	agtaagagaa	TTATGCAGTG	CTGCCATAAC	CATGAGTGAT	AACACTGCGG	3500
0	CCAACTTACT	TCTGACAACG	ATCGGAGGAC	CGAAGGAGCT	AACCGCTTTT	3550
15	TTGCACAACA	TGGGGGATCA	TGTAACTCGC	CTTGATCGTT	GGGAACCGGA	3600
. •	GCTGAATGAA	GCCATACCAA	ACGACGAGCG	TGACACCACG	ATGCCAGCAG	3650
20	CAATGGCAAC	AACGTTGCGC	AAACTATTAA	CTGGCGAACT	ACTTACTCTA	3700
	GCTTCCCGGC	: AACAATTAAT	AGACTGGATG	GAGGCGGATA	AAGTTGCAGG	3750
25	ACCACTTCTG	GCTCGGCCC	TTCCGGCTGG	CTGGTTTATI	GCTGATAAAT	3800
30	CTGGAGCCGG	G TGAGCGTGGG	TCTCGCGGTA	\ TCATTGCAGO	CACTGGGGCCA	3850
30	GATGGTAAGO	CCTCCCGTAT	CGTAGTTATC	TACACGACG	GGAGTCAGGC	3900
35	AACTATGGA:	r gaacgaaati	A GACAGATCG	TGAGATAGG	r GCCTCACTGA	3950
	TTAAGCATT	G GTAACTGTC	A GACCAAGTT	r actcatata	r actitagati	4000
40	GATTTAAAA	C TTCATTTT	A ATTTAAAAG	G ATCTAGGTG	A AGATCCTTT	4050
45	TGATAATCT	C ATGACCAAA	A TCCCTTAAC	G TGAGTTTTC	G TTCCACTGAC	3 4100
	CGTCAGACC	C CGTAGAAAA	G ATCAAAGGA	T CTTCTTGAG	A TCCTTTTTT	г 4150
50	CTGCGCGTA	A TCTGCTGCT	T GCAAACAAA	A <sub>.</sub> AAACCACCG	C TACCAGCGG	r 4200
	GGTTTGTTT	G CCGGATCAA	G AGCTACCAA	C TCTTTTTCC	G AAGGTAACT	G <b>42</b> 50
55	GCTTCAGCA	G AGCGCAGAT	A CCAAATACT	G TCCTTCTAG	T GTAGCCGTA	G 4300
60	TTAGGCCAC	C ACTTCAAGA	A CTCTGTAGO	A CCGCCTACA	T ACCTCGCTC	т 4350

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	GCTAATCCTG	TTACCAGTGG	CTGCTGCCAG	TGGCGATAAG	TCGTGTCTTA	4400
5	CCGGGTTGGA	CTCAAGACGA	TAGTTACCGG	ATAAGGCGCA	GCGGTCGGGC	4450
	TGAACGGGGG	GTTCGTGCAC	ACAGCCCAGC	TTGGAGCGAA	CGACCTACAC	4500
10	CGAACTGAGA	TACCTACAGC	GTGAGCATTG	AGAAAGCGCC	ACGCTTCCCG	4550
<b>4</b> F	AAGGGAGAAA	GGCGGACAGG	TATCCGGTAA	GCGGCAGGGT	CGGAACAGGA	4600
15	GAGCGCACGA	GGGAGCTTCC	AGGGGGAAAC	GCCTGGTATC	TTTATAGTCC	4650
20	TGTCGGGTTT	CGCCACCTCT	GACTTGAGCG	TCGATTTTTG	TGATGCTCGT	4700
	CAGGGGGGCG	GAGCCTATGG	AAAAACGCCA	GCAACGCGGC	CTTTTTACGG	4750
25	TTCCTGGCCT	TTTGCTGGCC	TTTTGCTCAC	ATGTTCTTTC	CTGCGTTATC	4800
••	CCCTGATTCT	GTGGATAACC	GTATTACCGC	CTTTGAGTGA	GCTGATACCG	4850
30	CTCGCCGCAG	CCGAACGACC	GAGCGCAGCG	AGTCAGTGAG	CGAGGAAGCG	4900
35	GAAGAGCGCC	CAATACGCAA	ACCGCCTCTC	CCCGCGCGTT	GGCCGATTCA	4950
	TTAATCCAGC	TGGCACGACA	GGTTTCCCGA	CTGGAAAGCG	GGCAGTGAGC	5000
40	GCAACGCAAT	TAATGTGAGT	TACCTCACTC	ATTAGGCACC	CCAGGCTTTA	5050
45	CACTTTATGC	TTCCGGCTCG	TATGTTGTGT	GGAATTGTGA	GCGGATAACA	5100
7.5	ATTTCACACA	GGAAACAGCT	ATGACCATGA	TTACGAATTA	A 5141	

- 50 (2) INFORMATION FOR SEQ ID NO:23:
  - (i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 21 bases

    - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
      - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

60 ATGTCNCCNG CNCCNCCNGC N 21

55

_	(2) INFORMATION FOR SEQ ID NO:24:		
5	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 21 bases</li><li>(B) TYPE: nucleic acid</li></ul>		•
10	<pre>(C) STRANDEDNESS: single (D) TOPOLOGY: linear</pre>		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	·	
15	ATGTCTCCAG CGCCGCCAGC G 21		
20	(2) INFORMATION FOR SEQ ID NO:25:  (i) SEQUENCE CHARACTERISTICS:		
	<ul><li>(A) LENGTH: 21 bases</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>		
25	(D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:		
30	ATGTCGCCTG CTCCACCTGC T 21		
35	(2) INFORMATION FOR SEQ ID NO:26:		
	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 21 bases</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>		
40	(D) TOPOLOGY: linear  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:		
45	ATGTCGCCAG CGCCACCAGC C 21		
50	(2) INFORMATION FOR SEQ ID NO:27:		
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 21 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> </ul>		
55	(D) TOPOLOGY: linear		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	•	
60	ATGTCCCCAG CCCCACCCGC A 21		

	(2) INFORMATION FOR SEQ ID NO:28:
5 10	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:
15	ATGTCGCCAG CGCCGCCAGC G 21
	(2) INFORMATION FOR SEQ ID NO:29:
20	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 25 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:
	Ser Pro Ala Pro Pro Ala Cys Asp Pro Arg Leu Leu Asn Lys Leu 1 5 10 15
30	Leu Arg Asp Asp His Val Leu His Gly Arg 20 25
	(2) INFORMATION FOR SEQ ID NO:30:
35	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 26 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:
	Ser Pro Ala Pro Pro Ala Cys Asp Pro Arg Leu Leu Asn Lys Leu 1 5 10 15
45	Leu Arg Asp Asp Xaa Val Leu His Gly Arg Leu 20 25 26
	(2) INFORMATION FOR SEQ ID NO:31:
50	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 25 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
55	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:
	Ser Pro Ala Pro Pro Ala Xaa Asp Pro Arg Leu Leu Asn Lys Leu 1 5 10 15
60	Leu Arg Asp Asp His Val Leu His Gly Arg

	(2) INFORMATION FOR SEQ ID NO:32:
5	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 14 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:  Xaa Pro Ala Pro Pro Ala Xaa Asp Pro Arg Leu Xaa Asn Lys  1 5 10 14
15	(2) INFORMATION FOR SEQ ID NO:33:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 9 amino acids
20	(R) TYPE: amino acid (D) TOPOLOGY: linear  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:
25	Pro Arg Leu Leu Asn Lys Leu Leu Arg 1 5 9
30	(2) INFORMATION FOR SEQ ID NO:34:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 45 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:
	GCCGTGAAGG ACGTGGTCGT CACGAAGCAG TTTATTTAGG AGTCG 45
40	(2) INFORMATION FOR SEQ ID NO:35:
45	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:
	CCNGCNCCNC CNGCNTGYGA 20
55	(2) INFORMATION FOR SEQ ID NO:36:
60	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 21 bases</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>

```
(D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:
 5
      NCCRTGNARN ACRTGRTCRT C 21
10 (2) INFORMATION FOR SEQ ID NO:37:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 69 bases
            (B) TYPE: nucleic acid
15
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:
20
      CCAGCGCCGC CAGCCTGTGA CCCCCGACTC CTAAATAAAC TGCCTCGTGA 50
      TGACCACGTT CAGCACGGC 69
25
     (2) INFORMATION FOR SEQ ID NO:38:
30
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 69 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
35
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:
      GGTCGCGGCG GTCGGACACT GGGGGCTGAG GATTTATTTG ACGGAGCACT 50
40
      ACTGGTGCAA GTCGTGCCG 69
45
     (2) INFORMATION FOR SEO ID NO:39:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 69 bases
            (B) TYPE: nucleic acid
50
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:
55
      CCAGCACCTC CGGCATGTGA CCCCCGACTC CTAAATAAAC TGCTTCGTGA 50
60
      CGACCACGTC CATCACGGC 69
```

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(2) INFORMATION FOR SEQ ID NO:40:
        (i) SEQUENCE CHARACTERISTICS:
5
            (A) LENGTH: 69 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:
      GGTCGTGGAG GCCGTACACT GGGGGCTGAG GATTTATTTG ACGAAGCACT 50
15
      GCTGGTGCAG GTAGTGCCG 69
20
     (2) INFORMATION FOR SEQ ID NO:41:
        (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 69 bases
             (B) TYPE: nucleic acid
25
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:
30
      CCAGCACCGC CGGCATGTGA CCCCCGACTC CTAAATAAAC TGCTTCGTGA 50
       CGATCATGTC TATCACGGT 69
35
      (2) INFORMATION FOR SEQ ID NO:42:
40
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 69 bases
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
 45
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:
       GGTCGTGGCG GCCGTACACT GGGGGCTGAG GATTTATTTG ACGAAGCACT 50
 50
       GCTAGTACAG ATAGTGCCA 69
 55
       (2) INFORMATION FOR SEQ ID NO:43:
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 37 bases
 60
              (B) TYPE: nucleic acid
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(C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:
 5
      GCTAGCTCTA GAAATTGCTC CTCGTGGTCA TGCTTCT 37
10
     (2) INFORMATION FOR SEQ ID NO:44:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 22 bases
15
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEO ID NO:44:
20
      CAGTCTGCCG TGAAGGACAT GG 22
25
     (2) INFORMATION FOR SEQ ID NO:45:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 24 bases
30
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:
35
      TGTGGACTTT AGCTTGGGAG AATG 24
40
     (2) INFORMATION FOR SEQ ID NO:46:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 22 bases
45
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:
50
      GGTCCAGGGA CCTGGAGGTT TG 22
55
      (2) INFORMATION FOR SEQ ID NO:47:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 31 bases
60
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
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	(D) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
5	ATCGATATCG ATAGCCAGAC ACCCCGGCCA G 31	. 🤸
10	(2) INFORMATION FOR SEQ ID NO:48:	:
15	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 36 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	
20	GCTAGCTCTA GACAGGGAAG GGAGCTGTAC ATGAGA 36	
25	(2) INFORMATION FOR SEQ ID NO:49:	
30	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 24 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:	
35	CTCCTTGGAA CCCAGGGCAG GACC 24	
40	(2) INFORMATION FOR SEQ ID NO:50:	
45	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 24 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:	
50	GGTCCTGCCC TGGGTTCCAA GGAG 24	
55	(2) INFORMATION FOR SEQ ID NO:51:	
60	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 27 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:
 5
      CTGCTCCGAG GAAAGGACTT CTGGATT 27
     (2) INFORMATION FOR SEQ ID NO:52:
10
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 27 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
15
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:
20
      AATCCAGAAG TCCTTTCCTC GGAGCAG 27
     (2) INFORMATION FOR SEQ ID NO:53:
25
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 28 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
30
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:
35
      CCCTCTGCGT CGCGGCGGCC CCACCCAC 28
     (2) INFORMATION FOR SEQ ID NO:54:
40
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 28 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
45
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEO ID NO:54:
50
      GTGGGTGGGG CCGCCGCGAC GCAGAGGG 28
     (2) INFORMATION FOR SEQ ID NO:55:
55
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 35 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
60
            (D) TOPOLOGY: linear
```

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:	••	
5	GACTCGAGGA TCCATCGATT TTTTTTTTTTT TTTTT 35		:
	(2) INFORMATION FOR SEQ ID NO:56:		
10	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 17 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>		•
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:		
20	GACTCGAGGA TCCATCG 17		
	(2) INFORMATION FOR SEQ ID NO:57:		
25	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 32 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>		
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:		
35	GCTAGCTCTA GAAGCCCGGC TCCTCCTGCC TG 32		
	(2) INFORMATION FOR SEQ ID NO:58:		
40	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 21 bases</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>		
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:		
50	CGAAATTAAC CCTCACTAAA G 21	•	
	(2) INFORMATION FOR SEQ ID NO:59:		
55	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 103 bases</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>		
60	(D) TOPOLOGY: linear		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:59:		

	TGCAGCAAGG GCTACTGCCA CACTCGAGCT GCGCAGATGC TAGCCTCAAG 50	)
5	ATGGCTGATC CAAATCGATT CCGCGGCAAA GATCTTCCGG TCCTGTAGAA 10	00
10	GCT 103	
	(2) INFORMATION FOR SEQ ID NO:60:	
15	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 103 bases  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	
20	(D) TOPOLOGY: linear  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:	
25	AGCTTCTACA GGACCGGAAG ATCTTTGCCG CGGAATCGAT TTGGATCAGC 50	)
	CATCTTGAGG CTAGCATCTG CGCAGCTCGA GTGTGGCAGT AGCCCTTGCT 10	00
30	GCA 103	
	(2) INFORMATION FOR SEQ ID NO:61:	
35	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 bases  (B) TYPE: nucleic acid	
40	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:	
45	TCTCGCTACC GTTTACAG 18	
	(2) INFORMATION FOR SEQ ID NO:62:	
50	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 25 bases  (B) TYPE: nucleic acid	
55	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	() GEOLIENGE DECORTEMION, CEO ID NO. 62	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:	

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(2) INFORMATION FOR SEQ ID NO:63:
        (i) SEQUENCE CHARACTERISTICS:
5
            (A) LENGTH: 18 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:
      GGGCCATGAC ACTGTCAA 18
15
     (2) INFORMATION FOR SEQ ID NO:64:
        (i) SEQUENCE CHARACTERISTICS:
20
             (A) LENGTH: 40 bases
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
25
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:64:
      GACCGCCACC GAGACCGCCT GGTGGGTACC TGTGGTCCTT 40
30
      (2) INFORMATION FOR SEQ ID NO:65:
         (i) SEQUENCE CHARACTERISTICS:
35
             (A) LENGTH: 32 bases
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
40
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:
       ATCGATATCG ATCAGCCAGA CACCCCGGCC AG 32
45
      (2) INFORMATION FOR SEQ ID NO:66:
         (i) SEQUENCE CHARACTERISTICS:
 50
             (A) LENGTH: 35 bases
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
              (D) TOPOLOGY: linear
 55
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:
       TCTAGATCTA GATCACCTGA CGCAGAGGGT GGACC 35
 60
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(2) INFORMATION FOR SEQ ID NO:67:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 33 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
10
    (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:
      AGTCGACGTC GACGTCGGCA GTGTCTGAGA ACC 33
15
     (2) INFORMATION FOR SEQ ID NO:68:
        (i) SEQUENCE CHARACTERISTICS:
20
            (A) LENGTH: 36 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
25
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
      AGTCGACGTC GACTCACCTG ACGCAGAGGG TGGACC 36
30
     (2) INFORMATION FOR SEQ ID NO:69:
        (i) SEQUENCE CHARACTERISTICS:
35
            (A) LENGTH: 62 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
40
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:
      CGCGTATGCC AGCCCGGCTC CTCCTGCTTG TGACCTCCGA GTCCTCAGTA 50
45
      AACTGCTTCG TG 62
50
   (2) INFORMATION FOR SEQ ID NO:70:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 61 bases
            (B) TYPE: nucleic acid
55
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:70:
60
      ATACGGTCGG GCCGAGGAGG ACGAACACTG GAGGCTCAGG AGTCATTTGA 50
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CGAAGCACTG A 61
5
     (2) INFORMATION FOR SEQ ID NO:71:
        (i) SEQUENCE CHARACTERISTICS:
10
            (A) LENGTH: 37 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:
15
      CTAGAATTAT GAAAAAGAAT ATCGCATTTC TTCTTAA 37
20
     (2) INFORMATION FOR SEQ ID NO:72:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 37 bases
25
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:
30
      TTAATACTTT TTCTTATAGC GTAAAGAAGA ATTGCGC 37
35
      (2) INFORMATION FOR SEQ ID NO:73:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 69 bases
40
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:
45
       CTAGAATTAT GAAAAAGAAT ATCGCATTTC ATCACCATCA CCATCACCAT 50
 50
       CACATCGAAG GTCGTAGCC 69
      (2) INFORMATION FOR SEQ ID NO:74:
 55
```

(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 62 bases(B) TYPE: nucleic acid(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

60

```
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:74:
 5
      TTAATACTTT TTCTTATAGC GTAAAGTAGT GGTAGTGGTA GTGGTAGTGT 50
      AGCTTCCAGC AT 62
10
     (2) INFORMATION FOR SEQ ID NO:75:
        (i) SEQUENCE CHARACTERISTICS:
15
            (A) LENGTH: 69 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
20
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:75:
      CTAGAATTAT GAAAAAGAAT ATCGCATTTC ATCACCATCA CCATCACCAT 50
25
      CACATCGAAC CACGTAGCC 69
30 (2) INFORMATION FOR SEQ ID NO:76:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 62 bases
            (B) TYPE: nucleic acid
35
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:76:
40
      TTAATACTTT TTCTTATAGC GTAAAGTAGT GGTAGTGGTA GTGGTAGTGT 50
      AGCTTGGTGC AT 62
45
     (2) INFORMATION FOR SEQ ID NO:77:
50
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 19 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
55
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77:
      TCCACCCTCT GCGTCAGGT 19
60
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60

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(2) INFORMATION FOR SEQ ID NO:78:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 18 bases
5
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:78:
10
      GGAGACGCAG TCCATCGA 18
15
     (2) INFORMATION FOR SEQ ID NO:79:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 62 bases
20
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:
25
      GCAGCAGTTC TAGAATTATG TCNCCNGCNC CNCCNGCNTG TGACCTCCGA 50
30
       GTTCTCAGTA AA 62
      (2) INFORMATION FOR SEQ ID NO:80:
35
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 49 bases
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
 40
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:
 45
       ACACTGGAGG CTCAAGAGTC ATTTGACGAA GCACTGAGGG TACAGGAAG 49
       (2) INFORMATION FOR SEQ ID NO:81:
 50
          (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 45 bases
              (B) TYPE: nucleic acid
              (C) STRANDEDNESS: single
 55
              (D) TOPOLOGY: linear
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:81:
```

CTAGAATTAT GAAAAAGAAT ATCGCATTTA TCGAAGGTCG TAGCC 45

```
(2) INFORMATION FOR SEQ ID NO:82:
5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 38 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
10
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:82:
15
     TTAATACTTT TTCTTATAGC GTAAATAGCT TCCAGCAT 38
     (2) INFORMATION FOR SEQ ID NO:83:
20
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 45 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
25
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:83:
      CTAGAATTAT GAAAAGAAT ATCGCATTTC TTCTTAAACG TAGCC 45
30
     (2) INFORMATION FOR SEQ ID NO:84:
35
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 38 bases
            (B) TYPE: nucleic acid
            (C) STRANDEDNESS: single
40
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:84:
45
      TTAATACTTT TTCTTATAGC GTAAAGAAGA ATTTGCAT 38
     (2) INFORMATION FOR SEQ ID NO:85:
50
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 25 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
55
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:85:
      Met Lys Lys Asn Ile Ala Phe Leu Leu Asn Ala Tyr Ala Ser Pro
        1 5
                                           10
60
```

```
Ala Pro Pro Ala Cys Cys Val Arg Arg Ala
                       20
     (2) INFORMATION FOR SEQ ID NO:86:
5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 31 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:86:
      Met Lys Lys Asn Ile Ala Phe His His His His His His His His
15
      Ile Glu Gly Arg Ser Pro Ala Pro Pro Ala Cys Cys Val Arg Arg
                                           25
      Ala
20
       31
     (2) INFORMATION FOR SEQ ID NO:87:
        (i) SEQUENCE CHARACTERISTICS:
25
            (A) LENGTH: 31 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:87:
30
      Met Lys Lys Asn Ile Ala Phe His His His His His His His
      Ile Glu Pro Arg Ser Pro Ala Pro Pro Ala Cys Cys Val Arg Arg
35
                       20
                                            25
                                                                30
      Ala
       31
40
      (2) INFORMATION FOR SEQ ID NO:88:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 7 amino acids
             (B) TYPE: amino acid
45
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:88:
       Thr Thr Ala His Lys Asp Pro
50
      (2) INFORMATION FOR SEQ ID NO:89:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
55
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:89:
```

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```
His Val Leu His
      1 4
     (2) INFORMATION FOR SEQ ID NO:90:
 5
        (i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 4 amino acids
           (B) TYPE: amino acid
           (D) TOPOLOGY: linear
10
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:90:
      Ser Arg Leu Ser
      1 4
15
     (2) INFORMATION FOR SEQ ID NO:91:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
20
           (B) TYPE: amino acid
           (D) TOPOLOGY: linear
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:91:
25
      Ser His Val Leu
      1 4
     (2) INFORMATION FOR SEQ ID NO:92:
30
        (i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
35
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:92:
      His Ser Arg Leu
40
     (2) INFORMATION FOR SEQ ID NO:93:
        (i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
           (D) TOPOLOGY: linear
45
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:93:
      Ala Val Asp Phe
50
      1 4
     (2) INFORMATION FOR SEQ ID NO:94:
        (i) SEQUENCE CHARACTERISTICS:
55
           (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:94:
60
```

```
Ser Leu Gly Glu
       1 4
     (2) INFORMATION FOR SEQ ID NO:95:
 5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:95:
      Ala Val Thr Leu
        1
15
     (2) INFORMATION FOR SEQ ID NO:96:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
20
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:96:
25
      Leu Leu Glu Gly
      (2) INFORMATION FOR SEQ ID NO:97:
30
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
35
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:97:
      Leu Ser Ser Leu
        1
40
      (2) INFORMATION FOR SEQ ID NO:98:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
45
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:98:
       Leu Gly Gln Leu
50
      (2) INFORMATION FOR SEQ ID NO:99:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 5 amino acids
55
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
```

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:99:

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```
Cys Xaa Leu Ser Ser
       (2) INFORMATION FOR SEQ ID NO:100:
  5
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
 10
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:100:
       Leu Leu Gly Gln
         1 4
 15
       (2) INFORMATION FOR SEO ID NO:101:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
 20
             (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:101:
 25
       Ser Ser Leu Leu
         1
                     4
       (2) INFORMATION FOR SEQ ID NO:102:
 30
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
 35
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:102:
       Gly Gln Leu Ser
40
       (2) INFORMATION FOR SEQ ID NO:103:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
 45
             (D) TOPOLOGY: linear
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:103:
       Cys Leu Ser Ser
 50
         1 4
       (2) INFORMATION FOR SEQ ID NO:104:
          (i) SEQUENCE CHARACTERISTICS:
 55
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO:104:
 60
```

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```
Leu Gln Ser Leu
       1 .4
     (2) INFORMATION FOR SEQ ID NO:105:
5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:105:
      Leu Gly Thr Gln
        1 4
15
     (2) INFORMATION FOR SEQ ID NO:106:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
20
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:106:
25
      Ala Leu Gln Ser
        1
     (2) INFORMATION FOR SEQ ID NO:107:
        (i) SEQUENCE CHARACTERISTICS:
30
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:107:
35
       Leu Leu Gly Thr
        1
      (2) INFORMATION FOR SEQ ID NO:108:
40
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
 45
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:108:
       Asn Ala Ile Phe
 50
      (2) INFORMATION FOR SEQ ID NO:109:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
 55
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
```

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:109:

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Leu Ser Phe Gln

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1 4
     (2) INFORMATION FOR SEQ ID NO:110:
 5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 22 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:110:
      Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val Leu Ser Lys Leu
                                     10 . 15
15
      Leu Arg Asp Ser His Val Leu
                      20
     (2) INFORMATION FOR SEQ ID NO:111:
20
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 24 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
25
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:111:
      His Ser Arg Leu Ser Gln Cys Pro Glu Val His Pro Leu Pro Thr
30
      Pro Val Leu Leu Pro Ala Val Asp Phe
                                    24 ·
                      20
     (2) INFORMATION FOR SEQ ID NO:112:
35
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 23 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
40
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:112:
      Ser Leu Gly Glu Trp Lys Thr Gln Met Glu Glu Thr Lys Ala Gln
                                          10
45
      Asp Ile Leu Gly Ala Val Thr Leu
                       20
     (2) INFORMATION FOR SEQ ID NO:113:
50
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 21 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
55
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:113:
      Leu Leu Glu Gly Val Met Ala Ala Arg Gly Gln Leu Gly Pro Thr
                      5
                                          10
60
```

```
Cys Leu Ser Ser Leu Leu
                  20 21
     (2) INFORMATION FOR SEQ ID NO:114:
5
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 16 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
10
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:114:
      Gly Gln Leu Ser Gly Gln Val Arg Leu Leu Gly Ala Leu Gln
                                          10
15
      Ser
       16
     (2) INFORMATION FOR SEQ ID NO:115:
20
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 22 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
25
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:115:
      Leu Leu Gly Thr Gln Leu Pro Pro Gln Gly Arg Thr Thr Ala His
                                         30
      Lys Asp Pro Asn Ala Ile Phe
                       20 22.
      (2) INFORMATION FOR SEQ ID NO:116:
35
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 25 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
40
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:116:
      Leu Ser Phe Gln His Leu Leu Arg Gly Lys Val Arg Phe Leu Met
45
      Leu Val Gly Gly Ser Thr Leu Cys Val Arg
                       20
      (2) INFORMATION FOR SEQ ID NO:117:
50
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
55
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:117:
       Met Pro Pro Ala
 60
```

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```
(2) INFORMATION FOR SEQ ID NO:118:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 5 amino acids
 5
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEO ID NO:118:
10
      Met Ala Pro Pro Ala
     (2) INFORMATION FOR SEQ ID NO:119:
15
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 6 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
20
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:119:
      Met Pro Ala Pro Pro Ala
25
     (2) INFORMATION FOR SEQ ID NO:120:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 7 amino acids
            (B) TYPE: amino acid
30
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:120:
      Met Ser Pro Ala Pro Pro Ala
35
                        5
     (2) INFORMATION FOR SEQ ID NO:121:
        (i) SEQUENCE CHARACTERISTICS:
40
             (A) LENGTH: 4 amino acids
             (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:121:
45
      Ala Pro Pro Ala
        1
     (2) INFORMATION FOR SEQ ID NO:122:
50
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 5 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
55
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:122:
      Pro Ala Pro Pro Ala
       1
60
```

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(2) INFORMATION FOR SEQ ID NO:123:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 6 amino acids
 5
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:123:
10
      Ser Pro Ala Pro Pro Ala
     (2) INFORMATION FOR SEQ ID NO:124:
15
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 4 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
20
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:124:
      Val Arg Arg Ala
       1 4
25
     (2) INFORMATION FOR SEQ ID NO:125:
        (i) SEOUENCE CHARACTERISTICS:
            (A) LENGTH: 5 amino acids
            (B) TYPE: amino acid
30
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:125:
      Val Arg Arg Ala Pro
35
     (2) INFORMATION FOR SEQ ID NO:126:
        (i) SEQUENCE CHARACTERISTICS:
40
            (A) LENGTH: 6 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:126:
45
      Val Arg Arg Ala Pro Pro
        1 5 6
      (2) INFORMATION FOR SEQ ID NO:127:
50
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 7 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
55
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:127:
      Val Arg Arg Ala Pro Pro Thr
                    5 7
60
```

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(2) INFORMATION FOR SEQ ID NO:128:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 8 amino acids
 5
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:128:
      Val Arg Arg Ala Pro Pro Thr Thr
10
                       5
     (2) INFORMATION FOR SEQ ID NO:129:
15
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 9 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
20
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:129:
      Val Arg Arg Ala Pro Pro Thr Thr Ala
                        5
     (2) INFORMATION FOR SEQ ID NO:130:
25
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 10 amino acids
            (B) TYPE: amino acid
30
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:130:
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val
35
                        5
     (2) INFORMATION FOR SEQ ID NO:131:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 11 amino acids
40
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:131:
45
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro
                                           10 11
        1
                        5
      (2) INFORMATION FOR SEQ ID NO:132:
50
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 12 amino acids
             (B) TYPE: amino acid
             (D) TOPOLOGY: linear
55
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:132:
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser
                                           10 12
             5
        1
60
```

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(2) INFORMATION FOR SEQ ID NO:133:
       (i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 13 amino acids
5
           (B) TYPE: amino acid
           (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:133:
     Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg
10
               5
     (2) INFORMATION FOR SEQ ID NO:134:
        (i) SEQUENCE CHARACTERISTICS:
15
            (A) LENGTH: 14 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:134:
20
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr
           5 10 14
25
     (2) INFORMATION FOR SEQ ID NO:135:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 15 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
30
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:135:
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
35
      1
                      5
                                        10
     (2) INFORMATION FOR SEQ ID NO:136:
        (i) SEQUENCE CHARACTERISTICS:
40
            (A) LENGTH: 16 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:136:
45
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
                                        10
50
      Leu
      (2) INFORMATION FOR SEQ ID NO:137:
         (i) SEQUENCE CHARACTERISTICS:
55
            (A) LENGTH: 17 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:137:
60
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Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
            5 10
      1
     Leu Val
 5
          17
     (2) INFORMATION FOR SEQ ID NO:138:
        (i) SEQUENCE CHARACTERISTICS:
10
            (A) LENGTH: 18 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:138:
15
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
                                         10
      Leu Val Leu
20
       18
     (2) INFORMATION FOR SEQ ID NO:139:
        (i) SEQUENCE CHARACTERISTICS:
25
            (A) LENGTH: 19 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:139:
30
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
      Leu Val Leu Thr
35
     (2) INFORMATION FOR SEQ ID NO:140:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 20 amino acids
40
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:140:
45
      Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser
        1
                      5
                                          10
                                                              15
      Leu Val Leu Thr Leu
50
     (2) INFORMATION FOR SEQ ID NO:141:
        (i) SEQUENCE CHARACTERISTICS:
55
            (A) LENGTH: 21 amino acids
            (B) TYPE: amino acid
            (D) TOPOLOGY: linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:141:
60
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	Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser  1 5 10 15
5	Leu Val Leu Thr Leu Asn 20 21
	(2) INFORMATION FOR SEQ ID NO:142:
1 0	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 22 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:142:
	Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser  1 5 10 15
20	Leu Val Leu Thr Leu Asn Glu 20 22
	(2) INFORMATION FOR SEQ ID NO:143:
25	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 23 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:143:
	Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser  1 5 10 15
35	Leu Val Leu Thr Leu Asn Glu Leu 20 23
	(2) INFORMATION FOR SEQ ID NO:144:
40	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 24 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:144:
<b>→</b> IJ	Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser 1 5 10 15
50	Leu Val Leu Thr Leu Asn Glu Leu Pro

## **CLAIMS**

We claim:

1. An isolated substantially homogeneous mpl ligand polypeptide.

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- 2. The mpl ligand polypeptide of Claim 1 selected from the group consisting of
  - (a) a fragment polypeptide;
  - (b) a variant polypeptide; and
  - (c) a chimeric polypeptide.

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- 3. The mpl ligand polypeptide of Claim 1 selected from the group consisting of
  - (a) the polypeptide that is isolated from a mammal;
  - (b) the polypeptide that is made by recombinant means; and
  - (c) the polypeptide that is made by synthetic means.

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- 4. The mpl ligand polypeptide of Claim 1 selected from the group consisting of
  - (a) the polypeptide that is human; and
  - (b) the polypeptide that is non-immunogenic in a human.
- 20 5. An isolated substantially homogeneous mpl agonist characterized in that:
  - (a) the agonist stimulates the incorporation of labeled nucleotides (<sup>3</sup>H-thymidine) into the DNA of IL-3 dependent Ba/F3 cells transfected with human *mpl* P; or
  - (b) the agonist stimulates <sup>35</sup>S incorporation into circulating platelets in a platelet rebound assay.
  - A fragment polypeptide according to Claim 2 represented by X-hTPO(7-151)-Y

where

WIIO

hTPO(7-151) represents the human TPO (hML) amino acid sequence from Cys<sup>7</sup> through Cys<sup>151</sup> inclusive;

X represents an amino group of Cys<sup>7</sup> or amino-terminus amino acid residue(s) selected from the group

Μ,

MA,

MPA,

MPPA, (SEQ ID NO: 117)

MAPPA, (SEQ ID NO: 118)

5

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MPAPPA, (SEQ ID NO: 119)

MSPAPPA, (SEQ ID NO: 120)

A,

PA,

PPA,

APPA, (SEQ ID NO: 121)

PAPPA, (SEQ ID NO: 122)
```

SPAPPA, (SEQ ID NO: 123)

Y represents the carboxy terminal group of Cys<sup>151</sup> or carboxy-terminus amino acid residue(s) selected from the group

٧. VR, VRR. VRRA, (SEQ ID NO: 124) VRRAP, (SEQ ID NO: 125) VRRAPP, (SEQ ID NO: 126) VRRAPPT, (SEQ ID NO: 127) VRRAPPTT, (SEQ ID NO: 128) VRRAPPTTA, (SEQ ID NO: 129) VRRAPPTTAV, (SEQ ID NO: 130) VRRAPPTTAVP, (SEQ ID NO: 131) VRRAPPTTAVPS, (SEQ ID NO: 132) VRRAPPTTAVPSR, (SEQ ID NO: 133) VRRAPPTTAVPSRT, (SEQ ID NO: 134) VRRAPPTTAVPSRTS, (SEQ ID NO: 135) VRRAPPTTAVPSRTSL, (SEQ ID NO: 136) VRRAPPTTAVPSRTSLV, (SEQ ID NO: 137) VRRAPPTTAVPSRTSLVL, (SEQ ID NO: 138) VRRAPPTTAVPSRTSLVLT, (SEQ ID NO: 139) VRRAPPTTAVPSRTSLVLTL, (SEQ ID NO: 140) VRRAPPTTAVPSRTSLVLTLN, (SEQ ID NO: 141) VRRAPPTTAVPSRTSLVLTLNE, (SEQ ID NO: 142) VRRAPPTTAVPSRTSLVLTLNEL, (SEQ ID NO: 143) VRRAPPTTAVPSRTSLVLTLNELP, (SEQ ID NO: 144),

amino-terminus amino acid residue extensions comprising one or more of the residues 176-332 of human ML as provided in Fig. 1 (SEQ ID NO: 1) and pegylated forms thereof.

7. A fragment polypeptide according to Claim 6 selected from the group TPO(1-153) and TPO(1-245).

5 8. A fragment polypeptide according to Claim 2, wherein the amino acid sequence of the fragment polypeptide comprises

SPAPPACDLRVLSKLLRDSHVL, (SEQ ID NO: 110)

HSRLSQCPEVHPLPTPVLLPAVDF,

10 (SEQ ID NO: 111)

SLGEWKTQMEETKAQDILGAVTL,

(SEQ ID NO: 112)

LLEGVMAARGQLGPTCLSSLL,

(SEQ ID NO: 113)

15 GQLSGQVRLLLGALQS,

(SEQ ID NO: 114)

LLGTQLPPQGRTTAHKDPNAIF,

(SEQ ID NO: 115)

LSFQHLLRGKVRFLMLVGGSTLCVR, and

20 (SEQ ID NO: 116)

combinations thereof.

- 9. The polypeptide of Claim 6 that is unglycosylated.
- 25 10. An isolated polypeptide encoded by a nucleic acid having a sequence that hybridizes under moderately stringent conditions to the nucleic acid molecules having a nucleic acid sequence provided in Fig. 1 (SEQ ID NO: 2).
  - 11. The polypeptide of Claim 11 that is biologically active.

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- 12. The polypeptide of Claim 1 selected from the group hML, hML<sub>153</sub>, hML(R153A, R154A), hML2, hML3, hML4, mML, mML2, mML3, pML, and pML2.
- 35 13. A polypeptide according to Claim 2, wherein the amino acid sequence of the polypeptide comprises amino acid residues 1 to X of Fig. 1 (SEQ ID NO: 1), where X is selected from the group 153, 155, 164, 174, 191, 205, 207, 217, 229, 245 and 332.

14. An isolated substantially homogeneous *mpl* ligand polypeptide sharing at least 80% sequence identity with the polypeptide of Claim 13.

- 5 15. The polypeptide of Claim 13 wherein X is 153.
  - 16. A chimera comprising the *mpl* ligand of Claim 13 fused to a heterologous polypeptide.
- 10 17. The chimera of Claim 16 wherein the heterologous polypeptide is an immunoglobin polypeptide.
  - 18. The chimera of Claim 16 wherein the heterologous polypeptide is an interleukin polypeptide.
- 15
  19. A chimera comprising the N-terminus residues 1 to about 153 to 157 of hML substituted with one or more, but not all, of the human EPO residues added or substituted into the N-terminus residues of hML at positions corresponding to the alignment shown in Fig. 10.
  - 20. An antibody that is capable of binding the mpl ligand polypeptide of Claim 13.
    - 21. A hybridoma cell line producing the antibody of Claim 20.

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- 25 22. An isolated nucleic acid molecule encoding the *mpl* ligand polypeptide of Claim 1.
  - 23. An isolated nucleic acid molecule encoding the *mpl* ligand polypeptide of Claim 13.
  - 24. An isolated nucleic acid molecule comprising the open reading frame nucleic acid sequence shown in Fig. 1 (SEQ ID NO: 2).
- The isolated nucleic acid molecule of Claim 24 encoding a *mpl* ligand polypeptide selected from the group hML, hML<sub>153</sub>, hML(R153A, R154A), hML<sub>2</sub>, hML<sub>3</sub>, hML<sub>4</sub>, mML<sub>4</sub>, mML<sub>2</sub>, mML<sub>3</sub>, pML, and pML<sub>2</sub>.

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26. An isolated nucleic acid molecule selected from the group consisting of

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- (a) a cDNA clone comprising the nucleotide sequence of the coding region of the mpl ligand gene;
- (b) a DNA sequence capable of hybridizing under stringent conditions to a clone of (a); and
- (c) a genetic variant of any of the DNA sequences of (a) and (b) which encodes a polypeptide possessing a biological property of a naturally occurring mpl ligand polypeptide.
- 10 27. An isolated DNA molecule having a sequence capable of hybridizing to a DNA sequence provided in Fig. 1 (SEQ ID NO: 2) under moderately stringent conditions, wherein the DNA molecule encodes a biologically active mpl ligand polypeptide.
- 15 28. The nucleic acid molecule of Claim 25 further comprising a promoter operably linked to the nucleic acid molecule.
  - 29. An expression vector comprising the nucleic acid sequence of Claim 25 operably linked to control sequences recognized by a host cell transformed with the vector.
    - 30. A host cell transformed with the vector of Claim 29.
- 31. A process of using a nucleic acid molecule encoding the *mpl* ligand polypeptide to effect production of the *mpl* ligand polypeptide comprising culturing the host cell of Claim 30.
  - 32. The process of Claim 31 wherein the *mpl* ligand polypeptide is recovered from the host cell.
  - 33. The process of Claim 31 wherein the *mpl* ligand polypeptide is recovered from the host cell culture medium.
- 34. A method of determining the presence of *mpl* ligand polypeptide, comprising hybridizing DNA encoding the *mpl* ligand polypeptide to a test sample nucleic acid and determining the presence of *mpl* ligand polypeptide DNA.

35. A method of amplifying a nucleic acid test sample comprising priming a nucleic acid polymerase reaction with nucleic acid encoding a *mpl* ligand polypeptide.

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- 36. A composition comprising the *mpl* ligand polypeptide of Claim 1 and a pharmaceutically acceptable carrier.
  - 37. A method for treating a mammal having or at risk for thrombocytopenia comprising administering to a mammal in need of such treatment a therapeutically effective amount of the composition of Claim 36.

The composition of Claim 36 further comprising a therapeutically effective amount of an agent selected from the group consisting of a cytokine, colony stimulating factor, and interleukin.

- 15 39. The composition of Claim 38 wherein the agent is selected from KL, LIF, G-CSF, GM-CSF, M-CSF, EPO, IL-1, IL-2, IL-3, IL-5, IL-6, IL-7, IL-8, IL-9 and IL-11.
- 40. A process for the biosynthesis of a human *mpl* ligand polypeptide, characterized by the amino acid sequence of one human *mpl* ligand polypeptide, such as that disclosed in Fig. 1 (SEQ ID NO: 1), comprising:
  - (a) growing a host cell culture containing a DNA isolate encoding the human *mpl* ligand polypeptide,
    - (b) recovering the human mpl ligand polypeptide from the culture, and
  - (c) purifying the *mpl* ligand polypeptide so as to obtain substantially homogeneous biologically active human *mpl* ligand polypeptide.

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:

1 tettectacceatetgetececagagggetgeetgetgtgeacttgggteetggageeettetecaceeggatagatteeteaeeettggeeegeettg

MetGluLeuThrGluLeuLeuValValMetLeuLeuLeuThrAlaArgLeuThrLeuSerSerProAlaProProAlaCysAsp

201 gacacceggecagaATGGAGCTGACTGAATTGCTCCTCGTGGTCATGCTTCTCCTAACTGCAAGGCTAACGCTGTCCAGCCCGGCTCCTCCTGCTTGTG

ACCTCCGAGTCCTCAGTAAACTGCTTCGTGACTCCCATGTCCTTCACAGCAGACTGAGCCAGTGCCCAGAGGTTCACCCTTTGCCTACACCTGTCCTGCT 301

GCCTGCTGTGGACTTTAGCTTGGGAGAATGGAAAACCCAGATGGAGGAGGAGCAAGGCACAGGACATTCTGGGAGCAGTGACCCTTCTGCAGGAGGAGTG ProAlaValAspPheSerLeuGlyGluTrpLysThrGlnMetGluGluThrLysAlaGlnAspIleLeuGlyAlaValThrLeuLeuLeuGluGlyVal

9

MetalaalaargglyginLeuGlyProThrCysLeuSerSerLeuLeuGlyGlnLeuSerGlyGlnValArgLeuLeuLeuGlyAlaLeuGlnSerLeuLe **ATGGCAGCACGCGGGACAACTGGGACCCACTTGCCTCTCATCCCTCCTGGGGCAGCTTTCTGGACAGGTCCGTCTCCTTCGGGGCCCTGCAGAGCCTCC** 501

GlyThrGlnLeuProProGlnGlyArgThrThrAlaHisLysAspProAsnAlaIlePheLeuSerPheGlnHisLeuLeuArgGlyLysValArgPhe TIGGAACCCAGCTICCTCCACAGGGCAGGACCACAGCTCACAAGGAICCCAAIGCCAICTICCIGAGCTICCAACACCTGCTCCGAGGAAAGGIGCGTIT 130 601

LeuMetLeuValGlyGlySerThrLeuCysValArgArgAlaProProThrThrAlaValProSerArgThrSerLeuValLeuThrLeuAsnGluLeu  ProasnargingserglyLeuLeugluthrannhethralaSeralaArgthrThrGlySerGlyLeuLeuLysTrpGlnGlnGlyPheArgAlaLysIle CCAAACAGGACTTCTGGATTGTTGGAGACAAACTTCACTGCCTCAGCCAGAACTACTGGCTCTGGGCTTCTGAAGTGGCAGCAGGGATTCAGAGCCAAGA 801

ProglyLeuLeu<u>AsnGlnThd</u>SerArgSerLeuAspGlnIleProGlyTyrLeuAsnArgIleHisGluLeuLeu<mark>AsnGlyThr</mark>ArgGlyLeuPhePro 220

TTCCTGGTCTGCTGAACCTCCAGGTCCCTGGACCAAATCCCCGGATACCTGAACAGGATACACGAACTCTTGAATGGAACTCGTGGACTCTTTCC 260

901

TGGACCCTCACGCAGGACCCTAGGAGCCCCGGACATTTCCTCAGGAACATCAGACACAGGCTCCCTGCCAACCTCCAGCCTGGATATTCTCCTTCC GlyProSerArgArgThrLeuGlyAlaProAspIleSerSerGlyThrSerAspThrGlySerLeuProProAsnLeuGlnProGlyTyrSerProSer

ProThrHisProProThrGlyGlnTyrThrLeuPheProLeuProProThrLeuProThrProValValGlnLeuHisPrcLeuLeuProAspProSerAla 

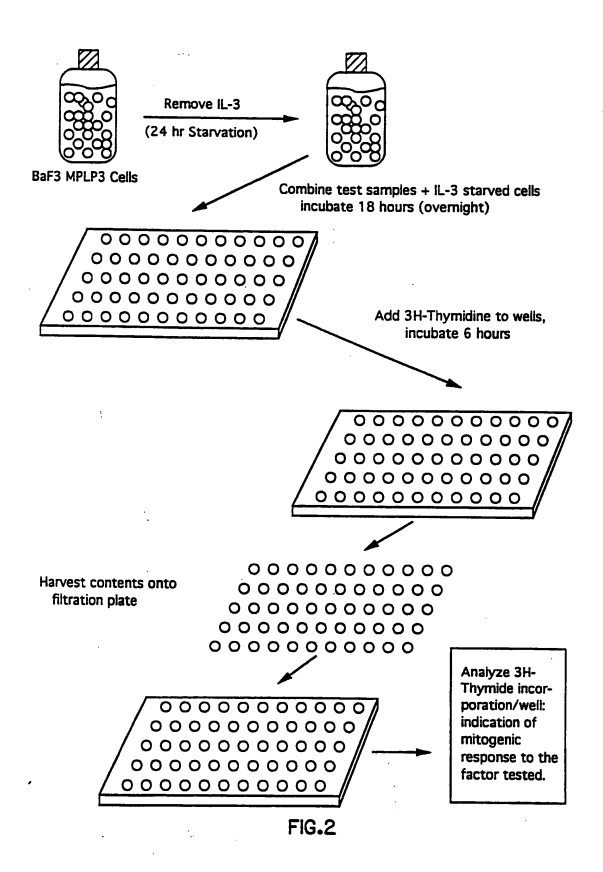
CTCCAACGCCCACCCCTACCAGCCCTCTTCTAAACACATCCTACACCCACAATTCTGTCTCAGGAAGGGTAAggttctcagacactgccgacatc ProThr ProThr ProThr Ser ProLeu Leu Asn Thr Ser Ilyr Thr His Ser Gln Asn Leu Ser Gln Glu Gly 330 320 310 1201 agcattgtctcatgtacagctcccttccctgcagggcgcccctggggagacaactggacaagatttcctacttctcctgaaacccaaagccctggtaaaa 1301

gggatacacaggactgaaaagggaatcatttttcactgtacattataaaccttcagaagctattttttaagctatcagcaatactcatcagagcagcta

1501

gaacagagggagagactaaccttgagtcagaaaacagagaaaagggtaatttcctttgcttcaaattcaaggccttccaacgccccatccctttactat 1601

FIG. I B



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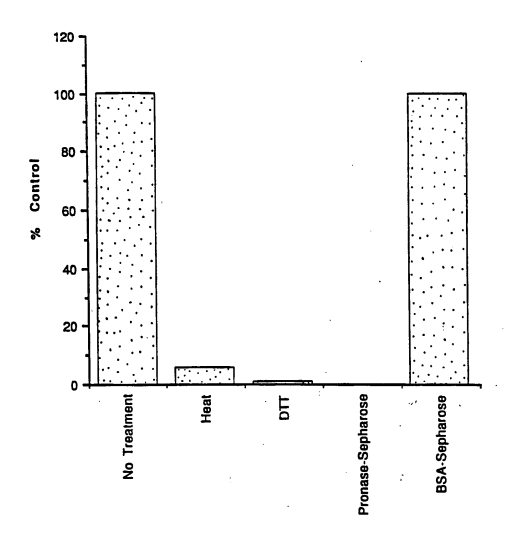


FIG.3

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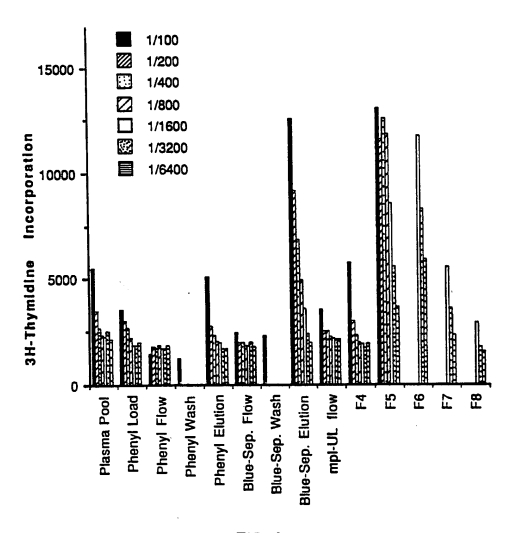


FIG.4

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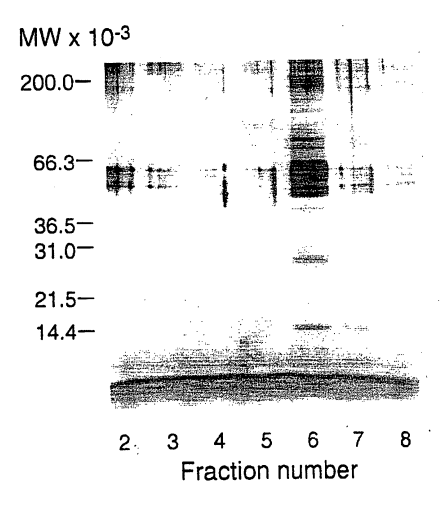


FIG. 5

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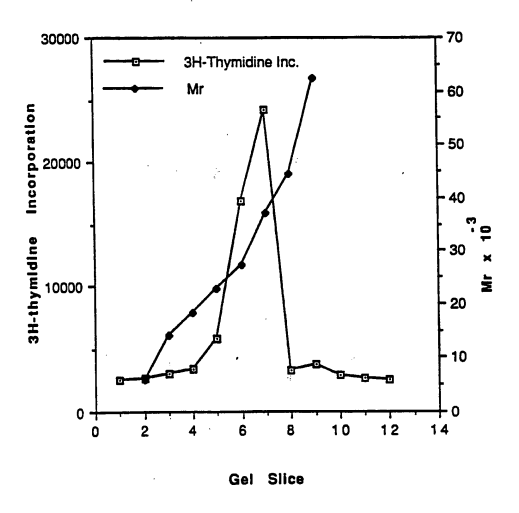


FIG.6

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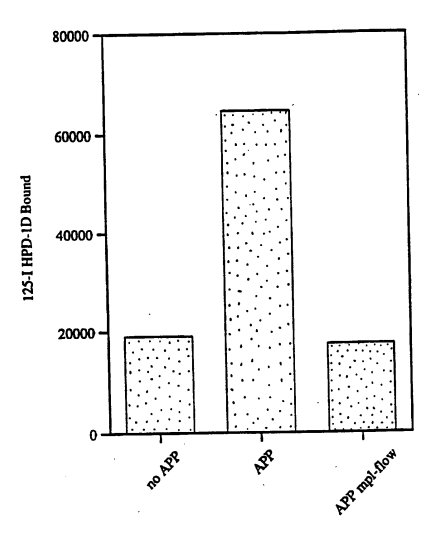


FIG.7

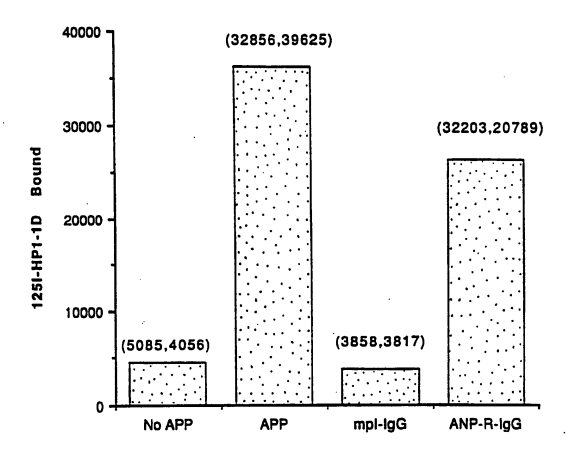


FIG.8

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1 GAATTCCTGG AATACCAGCT GACAATGATT TCCTCCTCAT CTTTCAACCT CACCTCTCT CATCTAAGAA TTGCTCCTCG TGGTCATGCT TCTCCTAACT CTTAAGGACC TTATGGTCGA CTGTTACTAA AGGAGGAGTA GAAAGTTGGA GTGGAGAGGA GTAGATTCTT AACGAGGAGC ACCAGTACGA AGAGGATTGA ר ר u > > L L L

10

-10

GAACTCCCAA CATTATCCCC TTTATCCGCG TAACTGGTAA GACACCCATA CTCCCAGGAA GACACCATCA CTTCCTCTAA CTCCTTGACC CAATGACTAT CTTGAGGGTT GTAATAGGGG AAATAGGCGC ATTGACCATT CTGTGGGTAT GAGGGTCCTT CTGTGGTAGT GAAGGAGATT GAGGAACTGG GTTACTGATA GGGCCGAGGA GGACGAACAC TGGAGGCTCA GGAGTCATTT GACGAAGCAC TGAGGGTACA GGAAGTGTCG TCTGACCACT CCTTCACAGC AGACTCGTGA LHS CTGCTTCGTG ACTCCCATGT SHS Ω œ ı ı A R L T L S S P A P P A C D L R V L S K GCAAGGCTAA CGCTGTCCAG CCCGGCTCCT CCTGCTTGTG ACCTCCGAGT CCTCAGTAAA GCGACAGGTC CGTTCCGATT 101

301 TCTTCCCATA TTGTCCCCAC CTACTGATCA CACTCTCTGA CAAGAATTAT TCTTCACAAT ACAGCCCGGA TTTAAAAAGCT CTCGTCTAGA AGAAGGGTAT AACAGGGTG GATGACTAGT GTGAGAGACT GTTCTTAATA AGAAGTGTTA TGTCGGGCGT AAATTFTCGA GAGCAGATCT

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**&** S <u>.</u> A ⋖  $\geq$ <u>ا</u> ه တ ⋖ 9 **5 2** FI  $\simeq$ م 6 1 LF > % **×** > **z**| **5** > -2 -LLE **& &** > H K F F HSRLSOCPEVHPLPTPVLLPENITVP 0 9 ш 9 ш H S ┢━. 9 0 R T T A H K D P N A I F L S F Q A A S A A P L R T I T A D T F R R T S · ш LS ے + ≥ œ ٥. Y L N SP = Z **\_**0 2 -0 -- MLVGGSTLCVRRAPPTTAVPSRTSLVLTLNELP GKLKLYTGEACRTGDR 1 C S S ٥ S SLDOIPGY SAPTPTPTSPLLNTSYTH FIG.10 S 2 ـ ۵. I L G A V T L L L E G V M A A R G O L G V W O G L A L L S E A V L R G O A L L V S 7 LOP  $\simeq$ 0 1 8 Z LPP 9 L L N ( - LG ALOS LLGIO - - - LPPOG SLTTLLRALGAOKEAISPPD SPAPPRLICDSRVLERYLLEAKEA S D T G X I P တ -⋖ 9 م  $\simeq$ S LHPLLPD u. S 9 -**5** 0 0 COMEETKAO C RMEVGOOAVE ۵. 0 3 ≪ LLK 1 L G TLPTPVVQ **&** 9 S 1 T G S X X 7 X 1- X · ~ ٥ 9 ∞ . ۵ 51 51 99 143 151 191 h-ML h-epo h-ML h-epo h-ML h-epo h-ML h-epo h-ML h-ML

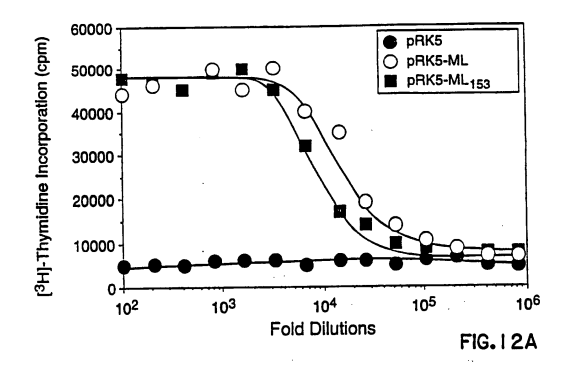
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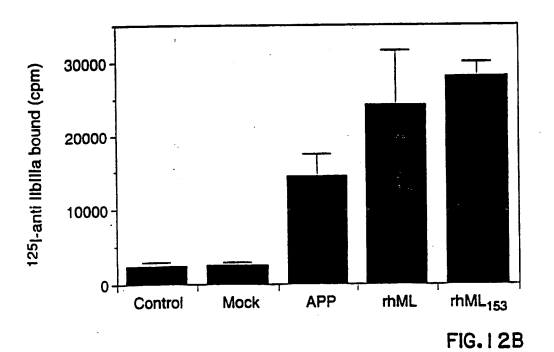
12/85

hML 201 hML2 197 hML3 179 hML4 175	QQGFRAKIPGLLNQTSRSLDQIPGYLNRIHELLNGTRGLFPGPSRRTLGA VPGPNPRIP EQDTRTLEWNSWTLSWTLTQDPRSPGHFLRNIRHRLPA VPGPNPRIP EQDTRTLEWNSWTLSWTLTQDPRSPGHFLRNIRHRLPA
251	PDISSGTSDTGSLPPNLQPGYSPSPTHPPTGQYTLFPLPPTL PDISSGTSDTGSLPPNLQPGYSPSPTHPPTGQYTLFPLPPTL
	TO
301	PLLPDPSAPTPTSPLLNTSYTHSQNLSQEG
297	PLLPDPSAPTPTSPLLNTSYTHSQNLSQEG
251	<del>-</del> -
	FIG. I 1B

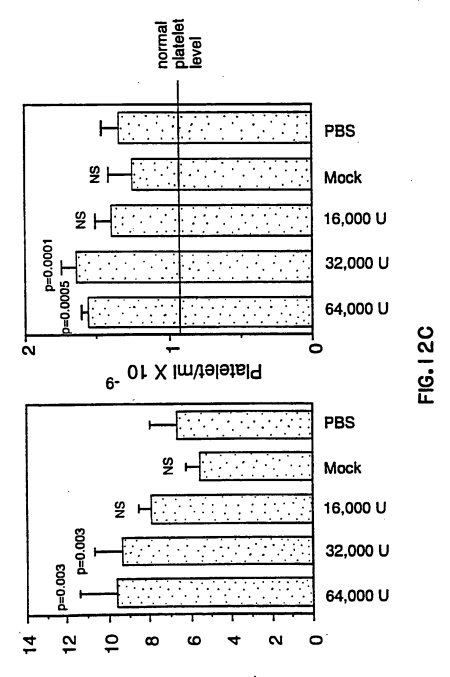
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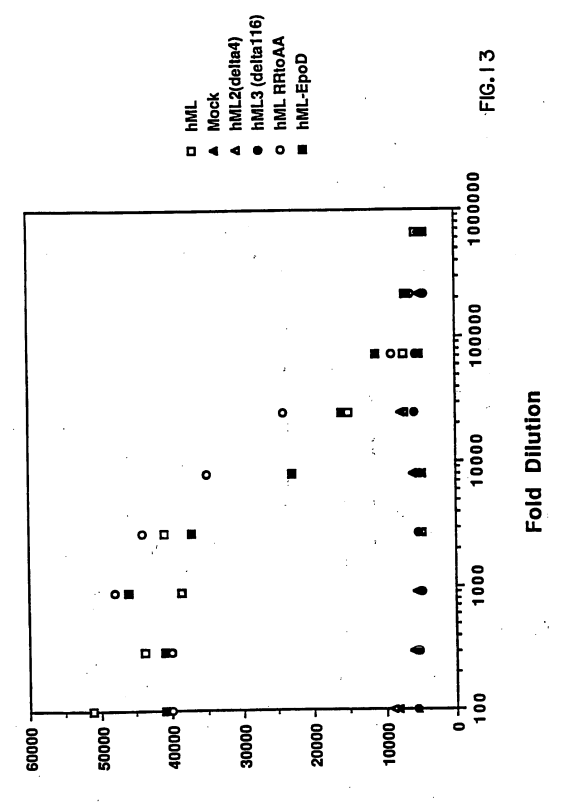


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 $^{\rm 835-S}$  Incorporation in Platelets X 10  $^{\rm 8}$ 

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[3H]-Thymidine Incorporation

16/85 SUBSTITUTE SHEET (RULE 26)

1 eccagectectttetettgttecetggteatgeetgeetecetgteteetgteteteeteeecacacacacacactateeteetagetateeetacaeee 201 acaggagagagctgaggaagttctgggggacaġggggatgatggtcaaggtcaggccaggaagccctgaggacagagactgtggggagactggggac 101 teetteetaatettgggagaeatetegtetggetggaegggaaaatteeaggatetaggeeacaetteteageagaeatgeecateettggggaggaggag 401 gggacacatgggcctggttattcctcttgtcacatgtggaacggtaggagatggaagacggagacagaacaagcaaaggagggccctggggcacagaggtc gacagtccactcaacccgtccaaaccctttccccataacacccataacagccataagagatttctctcatgtgggcaatatccgtgttcccacttcgaaagg gaaatttggatagccagggagtgaaaaccccaccaatcttaaacaagacctctgtgcttcttccccagcaacacaaatgtcctgccagattcctcctgga 701 801 1001

1301 ctggggggagggcaggcaactggaacctacaggcactgacctttgtcgagaagagtgtagccttcccagaatgggaggagcagggcagagcagggggtag 1501 aataagagagagctgcacttagggcttagcaaacacagtagtaagatggacacagccccaatccccattcttagctggtcattcctcgttagcttaag 1101 gaagtggeecaggeaggegtatgaectgetgtggaggggetgtgeeceaecgeeaeatgtetTCCTACCCATCTGCTCCCAGAGGGCTGCCTGCT 1201 TGCACTTGGGTCCTGGAGCCCTTCTCCACCCGgtgagtggccagcagggtgtggggttatgtgagggtagaaaggacagcaaagagaaatgggctcccag Start of cDNA sequence; Exon 1 >

FIG. 14A

1901 tcctgggcagctggtctcaggaagtcccagaactgttagcccatctcttggcctcagataatggagtatttcaggacttggagtccaaagaaaagc

agggggcagggaggcaggtgtgagctatgagacagatatgttagtgggcgcctaagacaaggtaagccctaaggtgggcatcacccagcaggtgcccgt

1701 teceaatattgaateaggtgeaageetetttgeacaaettgtgaaaggaggaggaageeatgtggggggteetgtgaaggaaeeggaaggggttetgeëa

2001 tccagtggctttatgtgtgggggtagatagggaaagaatagaggttaatttctcccataccgccttttaatcctgacctctagtggtcccagttacagct

gctggtcagcatctcaagccctccccagcatctgttcacctgccaggcagtctcttcctagaaacttggttaaatgttcactcttcttgctactttcag 2601 ccctgctagtttttttgtatttcgtagagccggggtttcaccatgttagtgaggctggtggcgaactcctgacctcaggtgatccaccgccttggaatc ccaaagtgctgggattacaggcatgagccactgcacccggcacaccatatgctttcatcacaagaaaatgtgagagaattcagggctttggcagttccag gtgcgatctcggctcaccacaaacctccgcctcccaggtacaagcgattctcctgtctcagcctcccaagtagcttggattacaggcatgagccacaca ttgtgcagttcccctccccagccccactccccaggaagttaccctcaacatattgcgcccgtttgccagttcctcacccaggccctgcatccat tttccactctcttctccaggctgaagccacaatactttccttctctatccccatcccagatttctctgacctaacaaccaaggttgctcagaatttaag 2301 gctaattaagatatgtgtatacatatcatgtcctgctgctctcagcaggggtaggtggcaccaaatccgtgtccgattcactgaggagtcctgacaaa

3001 GAGGCCCCAAACAGGGAGCCACGCCAGCCAGACACCCCCGGCCAGAATGGAGCTGACTGG tgagaacacacctgaggggctagggccatatggaaacatga MetGluLeuThrG

luLeuLeuLeu

3101 cagaaggggagagagaaaggagacacgctgcagggggcaggaagctgggggaacccattctcccaaaaataaggggtctgaggggtggattccctgggtt

FIG. 14B 3201 tcaggtctgggtcctgaatgggaattcctggaataccagctgacaatgatttcctcctcatcttcaacctcacctctctcatctaagAATTGCTCCTC

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2701

2201

ValValMetLeuLeuLeuThrAlaArgLeuThrLeuSerSerProAlaProProAlaCysAspLeuArgValLeuSerLysLeuLeuArgAspSerHisVal 3301 GIGGICAIGCITCICCIAACIGCAAGGCIAACGCIGICCAGCCCGGCICCICCIGCIIGIAACCICCICAGIAAACIGCIICGIGACICCAIG End of signal peptide^

43 LeuHisSerArgLeu

÷

:

3401 TCCTTCACAGCAGACTGgtgagaactcccaacattatcccctttatccgcgtaactggtaagacacccatactcccaggaagacaccatcacttcctcta

actecttgaeceaatgaetattetteecatattgteeceaectaetgateaeactetetgaeaagaattattetteaeaataeageeegetttaaaage 3501 3701 cagagccagrgcccagaggrrcacccrrrgccracccrgrccrgcrgcrgcrgcrgcrgrgagagcrrragcrrgggaaaacccagarggtaagaaagc  ${ t SerGInCysProGIuValHisProLeuProThrProValLeuLeuProAlaValAspPheSerLeuGlyGIuTrpLysThrGlnMet}$ 

1801 catccctaaccttggcttccctaagtcctgtcttcagtttcccactgcttcccatggattctccaacattcttgagctttttaaaaatatctcaccttca

gettggecacectaacecaatetacatteacetatgatgatgatageetgtggataagatgatggetggeaggtecaatatgtgaatagatttgaagetgaae 1001 accatgaaaagctggagagaaatcgctcatggccatgcctttgacctattccygttcagtcttcttaaattggcatgaagaagcaagactcatatgtcat 1101 ccacagatgacacaaagctgggaagtaccactaaaataacaaaagactgaatcaagattcaaaatcactgaaagactaggtcaaaaacaaggtgaaacaac

4301 agcagcctggccaacatggcgaaaccccgtctctactaagaatacaaaattagccgggcatggtagtgcatgcctgtaatcccagctacttggaaggctg

4401 aagcaggagaatcccttgaacccaggaggtggaggttgtagtgagctgagatcatgccaatgcactccagcctgggtgacaagagcaaaactccgtctca

4501 aaaagaaaaaaattetacatgtgtaaattaatgagtaaagteetatteeagettteaggeeacaatgeeetgetteeateatttaageetetggeeet

4601 agcacttcctacgaaaaggatctgagagaattaaattgcccccaaacttaccatgtaacattactgaagctgctattcttaaagctagtaattcttgtct

4701 gtttgatgtttagcatccccattgtggaaatgctcgtacagaactctattccgagtggactacacttaaatatactggcctgaacaccggacatccccct

4801 gaagacatatgctaattattaagagggaccatattaaactaacatgtgtctagaaagcagcagcctgaacagaaagagactagaagcatgttttatggg

FIG. 14C

=

5101 ccaggetggagtgcagtggcatgatetcaactcactgcaacetcaggeetcccggattcaagcgattetcetgcetcagtcteccaagtagetgggattac 5201 aggtgeceaceatgeceagetaatttttgtatttttggtagagatggggttteaceatgttggecaggetgatettgaaeteetgaeeteaggtgat

5401 cagaaagagtaaatttgcagcactagaaccaagaggtaaaagctgtaacagggcagatttcagcaacgtaagaaaaaaggagctcttctcactgaaacca

5601 accttggtcctgtccagttctcagcctgtatgattcactctgctggctactcctaaggctccccacccgctttagtgtgccctttgaggcagtgcgctt

GluGluThrLysAlaGlnAspIleLeuGlyAlaValThrLeuLeuLeuGluGlyValMetAlaAlaArgGlyGlnLeu 20/85

5801 GGACCCACTTGCCTCTCCTCCTGGGGCAGCTTTCTGGACAGGTCCGTCTCCTTGGGGCCCTGCAGAGCCTCCTTGGAACCCAGGtaagtccc GlyProThrCysLeuSerSerLeuLeuGlyGlnLeuSerGlyGlnValArgLeuLeuLeuGlyAlaLeuGlnSerLeuLeuGlyThrGln

5901 agtcaagggatctgtagaaactgttcttttctgactccagtcccactagaagacctgagggaagaagaagggctcttccagggagctcaagggcagaaggctg

6001 atctactaagagtgeteeetgeeageeacaatgeetgggtaetggeateetgtettteetaettagaeaagggaggeetgagatetggeeetggtgtttg

LeuProProGlnGlyArgThrThrAlaHisLysAspProAsnAlaIlePheLeuSerPheGlnHisLeuLeuArg

6101 GCCTCAGGACCATCCTCTGCCCTCAGGCTTCCTCCACAGGGCAGGACCACACCACAAGGATCCCAATGCCATCTTCCTGAGTTTCCAACACCTGCTCCG C for cDNA clone =========-^Alternative splice site GlyLysValArgPheLeuMetLeuValGlyGlySerThrLeuCysValArgArgAlaProProThrThrAlaValProSerArgThrSerLeuValLeu 6201 AGGAAAGGTGCGTTTCCTGATGCTTGTAGGAGGGTCCACCCTCTGCGTCAGGCGGGCCCCCACCACCACAGCTGTCCCCAGCAGAACCTCTTAGTCCTC ^End of EPO dormain

191 ThrLeuAsnGluLeuProAsnArgThrSerGlyLeuLeuGluThrAsnPheThrAlaSerAlaArgThrThrGlySerGlyLeuLeuLysTrpGlnGlnGly 6301 ACACTGAACGAGCTCCCAAACAGGACTTCTGGATTGTTGGAGACAAACTTCACTGCCTCAGCCAGAACTACTGGCTCTGGGCTTCTGAAGTGGCAGCAG

 ${ t PheArgAlaLysIleProGlyLeuLeuAsnGlnThrSerArgSerLeuAspGlnIleProGlyTyrLeuAsnArgIleHisGluLeuLeuAsnGlyThr}$ 6401 GATTCAGAGCCAAGATTCCTGGTCTGCTGAACCAAACCTCCAGGTCCCTGGACCAAATCCCCGGATACCTGAACAGGATACACGAACTCTTGAATGGAAC

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ArgGlyLeuPheProGlyProSerArgArgThrLeuGlyAlaProAspIleSerSerGlyThrSerAspThrGlySerLeuProProAsnLeuGlnPro TCGTGGACTCTTTCCTGGACCCTCACGCAGGACCCTAGGAGCCCCGGACATTTCCTCAGGAACATCAGACACAGGCTCCCTGCCACCAACCTCCAGCCT GlytyrSerProSerProThrHisProProThrGlyGlnTyrThrLeuPheProLeuProProThrLeuProThrProValValGlnLeuHisProLeuLeu 6601

5701 TTCCTGACCCTTCTGCTCCAACGCCCACCCCTACCAGCCCTCTTCTAAACACCATCCTACACCCACTCCCAGAATCTGTCTCTCAGGAAGGGTAAGGTTCTCA ProAspProSerAlaProThrProThrProThrSerProLeuLeuAsnThrSerTyrThrHisSerGlnAsnLeuSerGlnGluGly:STOP

6801 GACACTGCCGACATCAGCATTGTCTCATGTACAGCTCCCTTCCCTGCAGGCGCCCCCTGGGAGACAACTGGACAAGATTTCCTACTTTCTCCTGAAACCC

AAAGCCCTGGTAAAAGGGATACACAGGACTGAAAAGGGAATCATTTTTCACTGTACATTATAAAACCTTCAGAAGCTATTTTTAAGCTATCAGAATAC

1001 TCATCAGAGCAGCTAGCTCTTTGGTCTATTTTCTGCAGAAATTTGCAACTCACTGATTCTCTACATGCTCTTTTTTCTGTGATAACTCTGCAAAGGCCTGG 7101 GCTGGCCTGGCAGTTGAACAGAGGGAGAGACTAACCTTGAGTCAGAAAACAGAGAAAGGGTAATTTCCTTTGCTTCAAATTCAAGGCCTTCCAACGCCCC 7301 ctatacactagacaaaactgagcctgtataaggaataaatgggagcgccgaaaagctccctaaaaagcaagggaaaggtgttcttcgagggtggcaatag

CATCCCCTTTACTATCATTCTCAGTGGGACTCTGATCCCATATTCTTAACAGATCTTTACTCTTGAGAAATGAATAAGCTTTCTCTCAGAAAtgctgtcc

7401 atcccctcaccctgccacccaaacaaaaagctaacaggaagccttggagagcctcacaccccaggtaaggctgtgtagacagttcagtaaagacagg

7701 aaatgcatctaaaaagcagctctgtgtgtgaccaccataaactctgctaggggatctctaaaaaaggagtcaggcttatgggggctttgcaaataagtgctgcc

7601 gaagtgcaggaagtggcatgagtcaggctccttgagctcacacagcaggagaacaagtacaagtcaagtacaagttgaaggctcatttcccagttcccgc

7801 ttggtgctcaggaaaaggtttgtgttgcacaaaacacaaattccactgc

FIG. 14E

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	—66.3 —55.4	
	36.5 31.1	

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FIG. 15

-21.5

FIG. I 6A

1 GAGTECTING CECACETETE INCEACEEGA CICINGEGAA AGAAGEACAG AAGETEAAGE EGEETECAIG GECECAGGAA AGAITEAGGG GAGAGGEEEE uThrLeuSer alalaArgLe LeuLeuAlaV Met GluLeuThrA spLeuLeuLe uAlaAlaMet

:

;

101 ATACAGGGAG CCACTTCAGT TAGACACCCT GGCCAGAATG GAGCTGACTG ATTTGCTCCT GGCGGCCATG CTTCTTGCAG TGGCAAGACT AACTCTGTCC 20 10

rGlnCysPro AspValAspPro euGlyAlaVal GCCGACTGAG TCAGTGTCCC GACGTCGACC CITITGICIAT CCCIGITICIG CIGCCIGCIG IGGACITITAG CCIGGGAGAA IGGAAAACCC AGACGGAACA GAGCAAGGCA CAGGACAITC IAGGGGCAGI GlnAspileL Serprovala laproalacy saspproarg Leuleuasni ysleuleuar gaspserhis Leuleuhiss erargleuse agcecegtag etecteceng traceceana etectada nSerLysAla 9 LeuSeril eProValLeu LeuProAlaV alAspPheSe rLeuGlyGlu TrpLysThrG lnThrGluGl 201

LeuglyAlai euglnglyle uleuglyThr ginglyArgt hrThrAlaHi siysAspPro AsnAlaieup heleuSerle uglnglnleu LeuArgGlyLys Troggogoco rocaggocot cotaggaaco cagggaagaa coacagotoa caaggaococo aatgocotot rotroaggoto goaacaacto citogggaa TCGCCTCCTC CCTGCCTCTC ATCCCTCCTG GGACAGCTTT CTGGGCAGGT 120 GTCCCTTCTA CTGGAGGGG TGATGGCAGC ACGAGGACAG TTGGAACCCT 110 401 501

Leugluglyv almetalaal aargglygin LeugluPros erCysleuSe rSerLeuLeu GlyginLeus erGlyginVa largLeuLeu

80

Valargph eLeuLeuLeu ValgluglyP roThrLeuCy sValargarg ThrLeuProT hrThrAlaVa 1ProSerSer ThrSerGlnL euLeuThrLeu 601 AGGRGGGCTT CCTGCTTCTG GTAGAAGGTC CCACCCTCTG TGTCAGACGG ACCTGCCAA CCACAGGTGT CCTAAGCAGT ACTTCTCAAC TCCTCAACA 150

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FIG. 1 6B

190 170 AsnLysPhe ProAsnArgT hrSerGlyLe uLeuGluThr AsnPheSerV alThrAlaAr gThrAlaGly ProGlyLeuL euSerArgLe uGlnGlyPhe 701 AAACAAGTTC CCAÄÄCAGGA CTTCTGGATT GTTGGAGACG ĀĀCTTCĀGTG TCACAGCCAG AACTGCTGGC CCTGGACTTC TGAGCAGGCT TCAGGGATTC

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	230 GINThrSerA rgSerProva 1G1n11eSer G1yTyrLeuA snArgThrHi sG1yProval AsnG1yThrHis CAAACCTCCA GGTCCCCAGT CCAAATCTCT GGATACCTG <del>A ACAGGACA</del> CA CGGACCTGTG AATGGAACTC	260 hrLeuGluAl aSerAspIle SerProGlyA laPheAsnLy sGlySerLeu AlaPheAsnL euGlnGlyGly CCCTGGAAGC CTCAGACATC TCGCCCGGAG CTTTCAACAA AGGCTCCCTG GCATTCAACC TCCAGGGTGG	300 erProProGl nLeuHisPro CTCCACCCCA GCTCCACCCC	Thr ACATAGCGCG	TCTCGGGGAC AAGCTTCCCC AGGAAGGCTG AGAGGCAGCT GCATCTGCTC CAGATGTTCT GCTTTCACCT
4	230 GINThrSerA rgSerProva iginileser GlyTyrLeuA snArgThrHi sglyProval CAAACCrcca GGTCCCCAGT CCAATCTCT GGATACCTGA ACAGGACACA CGGACCTGTG	260 SerProGlyA laPheAsnLy sGlySerLeu AlaPheAsnL euGlnGlyGly TCGCCCGGAG CTTTCAACAA AGGCTCCCTG GCATTCAACC TCCAGGGTGG	290 pGlyHisThr ProPheProP roSerProAl aLeuProThr ThrHisGlyS erProProGl nLeuHisPro rggacacaca cccrrcccrc crrcaccrc crrcccacc acccargar crccaccca gcrccaccc	330 AsnSerThrA laProHisPr oValThrMet TyrProHisP roArgAsnLe uSerGlnGlu Thr AACTCTACCG CCCTCATCC AGTCACAATG TACCCTCATC CCAGGAATTT GTCTCAGGAA ACATAGCGCG	CAGATGTTCT
	snargThrHi <u>Acagaca</u> ca	260 sGlySerLeu AGGCTCCCTG	ThrHisGlyS ACCCATGGAT	roArgAsnLe CCAGGAATTT	GCATCTGCTC
	GlytyrLeua Ggatacctgā	<b>laPheAsnLy</b> CTTTCAACAA	290 Propheprop roserproal aleuprothr ThrHisGlyS CCCTTCCCTC CTTCCCTGC CTTGCCCACC ACCCATGGAT	Tyr ProHisP TACCCTCATC	AGAGGCAGCT
	220 1GlnIleSer CCAAATCTCT	SerProGlyA TCGCCCGGAG	roSerProAl CTTCACCTGC	320 AsnSerThrA laProHisPr oValThrWet AACTCTACCG CCCTCATCC AGTCACAATG	AGGAAGGCTG
	rgSerProVa GGTCCCCAGT	250 aSerAspIle CTCAGACAIC	ProPheProP CCCTTCCCTC	laProHisPr CCCCTCATCC	AAGCTTCCCC
			280 pglyhisthr tggacacaca		
	210 yGlnLeuAsn TCAGCTAAAT	SerLeuGlnT TCACTTCAGA	euAlaProAs TTGCTCCTGA	310 rThrMetPro CACCATGCCT	CTGCAGCTTC
	210 ArgValLysi leThrProgl yGlnLeuAsn AGAGTCAAGA TTACTCCTGG TCAGCTAĀĀĀ	240 GlyLeuPh eAlaGlyThr SerLeuGlnT GGGCTCTT TGCTGGAACC TCACTTCAGA	270 Leupropro SerproSerL euAlaProAs CTTCCTCCT TCTCCAAGCC TTGCTCCTGA	310 Leupheproa spproSerTh rThrMetPro CTGTTTCCTG ACCCTTCCAC CACCATGCCT	CAGTGAGCGT
210000000000000000000000000000000000000	210 ArgValLysi leThrProGl yGlnLeuAsn 801 AGAGTCAAGA TTACTCCTGG TCAGCTAAAT	240 GlyLeuPh eAlaGlyThr SerLeuGlnT	270 LeuProPro SerProSerL euAlaProAs 1001 ACTCCTCCT TCTCCAAGCC TTGCTCCTGA	310 LeuPheProA spProSerTh rThrMetPro 1101 CTGTTTCCTG ACCCTTCCAC CACCATGCCT	1201 GGCACTGGCC CAGTGAGCGT CTGCAGCTTC
1	801	901	1001	1101	1201

1401 TCTTTGGTCT ATTTTCGGTA TAAATTTGAA AATCACTAAT TCT

1301 AAAAGGCCCT GGGGAAGGGA TACACAGCAC TGGAGATTGT AAAATTTTAG GAGCTATTT TITTTAACCT ATCAGCAATA TTCATCAGAG CAGCTAGCGA

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MetGluLeuThrAspLeuLeuAlaAlaMetLeuLeuAlaAlaMetLeuLeuAlaValAlaAlaAlaAlaBetLeuLeuLeuAlaValAlaArgLeuThrLeuSer atacagggagccacttcagttagacacctggccagaATGGAGCTGACTGATTTGCTCCTGGCGGCCATGCTTCTTGCAGTGGCAAGACTAACTCTGTCC

Ser ProValAla ProAlaCys Asp ProArg Leu Leu Asn Lys Leu Leu Arg Asp Ser His Leu Leu His Ser Arg Leu Ser Gln Cys ProAsp Val Asp Pro 201 LeuSerIleProValLeuLeuProAlaValAspPheSerLeuGlyGluTrpLysThrGlnThrGluGlnSerLysAlaGlnAspIleLeuGlyAlaVal CTTTGTCTATCCCTGTTCTGCTGCTGCTGTGGACTTTAGCCTGGGAGAATGGAAAACCCAGACGGAACAGAGCAAGGCACAGGACATTCTAGGGGCAGG 301

SerLeuLeuLeuGluGlyValMetAlaAlaArgGlyGlnLeuGluProSerCysLeuSerSerLeuLeuGlyGlnLeuSerGlyGlnValArgLeuLeu GTCCCTTCTACTGGAGGGÄGTGATGGCAGGÄGGÄGAGTTGGAACCCTCCTĞCCTCTCTATCCCTCGGÄCAGCTTCTGGĞCAGGTTCGČCTCCTC 80 401

LeuglyAlaLeuglnGlyLeuLeuglyThrGlnLeuProLeuglnGlyArgThrThrAlaHisLysAspProAsnAlaLeuPheLeuSerLeuGlnGlnLeu TrggggggcccrgcAgggccrccTAggAACC<u>CAGCTTCCTA</u>CAGGGCAGGACCACAGCTCACAAGGACCCCAATGCCCTCTTCTTGAGCTTGCAACAAC 110 501 LeuArgGlyLysValArgPheLeuLeuLeuValGluGlyProThrLeuCysValArgArgThrLeuProThrThrAlaValProSerSerThrSerGln TGCTTCGGGGÄAÄGGTGCGCTTCCTGCTTGTAGAAGGTCCCACCCTCTGTGTCAGÄCGGACCCTGCCAACCACAGCTGTCCCAAGCAGTACTTCTCA 601

160

LeuLeuThrLeuAsnLysPhePro<u>AsnArgThI</u>SerGlyLeuLeuGluThr<u>AsnPheSer</u>ValThrAlaArgThrAlaGlyProGlyLeuSerArg ACTCCTCACACTAAACAAGTTCCCAAACAGGACTTCTGGATTGTTGGAGACTTCAGTGTCACAGACCAGAACTGCTGGCCTGGACTTCTGAGCAGG

FIG. 17A

LeuGlnGlyPheArgValLysIleThrFroGlyGlnLeuAsnGInThTSerArgSerFroValGlnIleSerGlyTyrLeuAsnArgThTHisGlyProVal CTTCAGGGATTCAGAGTCAAGATTACTCCTGGTCAGCTAAATCAAACCTCCAGTCCCAGTCCAAATCTGGATACCTGAACAGGACACAGGACCTG

250

ASIGIYTHI ISGIY LeuPheAlaGlyThr Ser LeuGlnThr LeuGluAla Ser Aspile Ser ProGlyAla PheAsn LysGlySer LeuAla PheAsn TGAATGGAACT CATGGGCT CTTTGCTGGAACCT CAGACCCTGGAAGCCT CAGACATCT CGCCCGGAGCTTT CAACAAAGCCT CCTGGCATT CAA

LeuGlnGlyGlyLeuProProSerProSerLeuAlaProAspGlyHisThrProPheProProSerProAlaLeuProThrThrHisGlySerProPro cctccAgggTggActTcctccTtccCAAgccTTgCTccTgATgGACACACACCCTTcCCTTCACCTGCCTTGCCCACCACCACCATGGATCTCCACC

GlnLeuHisProLeuPheProAspProSerThrThrMetProAsnSerThHAlaProHisProValThrMetTyrProHisProArgAshLeuSerGlnGlu CAGCTCCACCCCTGTTTCCTGACCCTTCCACCACCATGCCTAACTCTACCGCCCTCATCCAGTCAATGTACCCTCATCCAGGAATTTGTCTCAGG 1101

AAACATAGegegggeactggeccagtgagegtctgcagetteteteteggggacaagetteeccaggaaggetgagaggeagetgcatetgetecagatgtt  1401

1501 вааааааааааааааааааааааааааааааааа

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hML3	1 SPAPPACDL RVLSKLLRDSHVLHSRLSQCPEVHPLPTPVLLPAVDFSLGE
mML3	1 SPVAPACDPRLLNKLLRDSHLLHSRLSQCPDVDPLS 1 PVLLPAVDFSLGE
hML3	51 WKTOMEETKAODILGAVTLLLEGVMAARGOLGPTCLSSLLGOLSGOVRLL
mML3	51 WKTOTEOSKAODILGAVSLLLEGVMAARGOLEPSCLSSLLGOLSGOVRLL
hML3	101 LGALQSLLGTQLPPQGRTTAHKDPNAIFLSFQHLLRGKDFWIVGDKLHCL
mML3	101 LGALQGLLGTQLPLQGRTTAHKDPNALFLSLQQLLRGKDFWIVGDELQCH
hML3 mML3	151 S Q N Y WL WAS EVAAGIQSQD. S WS AEPNLQVPGPNPRIPEQDTRTLEWNS W 151 S Q N C WP WT S E Q AS GIQSQD Y S WS AKS N L Q V P S P N L W I P E Q D T R T C E WN S W
hML3	200 TLSWTLTODPRSPGHFLRNIRHRLPATOPPAWIFSFPNPSSYWTVYALPS
mML3	201 ALCWNLTSDPGSLRHLARSFOORLPGIOPPGWTSSFSKPCS
ьмг3	FIG. 18

L L P A V D F S L G E	L G O L S G O V R L L	RFLLLVEGPTU	A R T A G P G L L S R	LFAGISLOTIE	S P A L P I T H G S P	FIG. 19
L L P A V D F T L G E	L V O L S G O V R L L	RFLLLVVGPSU	A R T T G S G F L K W	LFPGPPGALG	S P T S P S P T	
L L P A V D F S L G E	L G O L S G O V R L L	RFLMLVGGSTL	A R T T G S G L L K W	LFPGPSRRTLG	P P T L P T P V	
PVAPACDPRLLNKLLRDSHLLHSRLSOCPDVDPLSIPVL	K T O T E	LGALOGLLGTOLPLOGRITAHKDPNALFLSLOOLLRGKVR	CVRRILLPITAVPSSTSOLLTLNKFPNERGISGLLETNERSVTA	RVKIITPGOLNOOMSRSPVOISGYLNIRTHGPVNGTHG	ASD I SPGAFN KG SLAFN LOGG LPPSPSLAPDGH-TPFPPS	POLHPLEPDPSTTMPNSSEAPHPVIMYPHPRNLS10ETVOLOPLLPDPSAITPNSSEEVOLDFAAHPHFONSSOEEVOLHPLLPDPSAPTPTPTSPLLRESYTHSONESOEG
PAPPACDPRLLNKLLRDSHVLHGRLSOCPDINPLSTPVL	K T O T E	LGALODLLGMOLPPOGRITAHKDPSAIFLNFOOLLRGKVR	CAKRAPPAIAVPSSTSPFHTLNKLPNERGESGLLETNESSVTA	RAKI-PGLLNOOMSRSLDOIPGHONGTHGPLSGIHG	APDIPPATSGMG SRPTYLOPGESPSPAHPSPGRYTLFSPS	
PAPPACDLRVLSKLLRDSHVLHSRLSOCPEVHPLPIPVL	K T O ME	LGALOSLLGTOLPPOGRITAHKDPNAIFLSFOHLLRGKVR	CVRRAPPTTAVPSRTSLVLTLNELPNERGISGLLETNEFTASA	RAKI-PGLLNOOMSRSLDOIPGYLNRIHELLNGTRG	APDISSGISD TG SLPPNLOPGYSPSPTHPPTGOYTLFPLP	
1 1 1 N N N	Z Z Z	101	151	201	251	300 297
m-ML o-ML	m-ML p-ML h-ML	m-ML p-ML h-ML	m-ML p-ML h-ML	m-ML p-ML h-ML	m-ML p-ML h-ML	m-ML p-ML h-ML

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Ser Pro Ala Pro Pro Ala Cys Asp Pro Arg Leu Leu Asn Lys Leu Leu Arg Asp Ser His Val Leu His Gly Arg Leu Ser Gln Cys Pro Asp Ile Asn Pro 

LeuSerThrProValLeuLeuProAlaValAspPheThrLeuGlyGluTrpLysThrGlnThrGluGlnThrLysAlaGlnAspValLeuGlyAlaThr9

ThrLeuLeuLeuGluAlaValMetThrAlaArgGlyGlnValGlyProProCysLeuSerSerLeuLeuValGlnLeuSerGlyGlnValArgLeuLeu AACCCTTCTGCTGGAGGCAGTGATGACAGCACGGGGACAAGTGGGACCCCCTTGCCTTCATCCCTGCTGGTGCAGCTTTCTGGACAGGTTCGCCTCCTC 80 201

LeuGlyAlaLeuGlnAspLeuLeuGlyMetGlnLeuProProGlnGlyArgThrThrAlaHisLysAspProSerAlaIlePheLeuAsnPheGlnGlnLeu CTCGGGGCCCTGCAGGACCTCCTTGGAATGCAGCTTCCTCCACAGGGAAGGACCACAGGTCACAAGGATCCCAGTGCCATCTTCCTGAACTTCCAACAAC 110 301

120

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LeuGlnAlaPheArgAlaLysIleProGlyLeuLeuAsnGlnThrSerArgSerLeuAspGlnIleProGlyHisGlnAsnGlyThrHisGlyProLeuSer CTGCAGGCATTCAGAGCCAAGATTCCTGGTCTGCTGAACCAAACCTCCAGGTCCCTAGACCAAATCCCTGGACACCAGAATGGGACACACGGACCCTTGA 330

GlylleHisGlyLeuPheProGlyProGlnProGlyAlaLeuGlyAlaProAspIleProProAlaThrSerGlyMetGlySerArgProThrTyrLeu 901 CCTCTGCTTCCTGACCCCTCTGCGATCACACCCCAACTCTACCAGTCCTCTTTTTTGCAGCTCACCTCATTTCCAGAACCTGTCTCAGGAAGAAGTAAG GlnProGlyGluSerProSerProAlaHisProSerProGlyArgTyrThrLeuPheSerProSerProThrSerProSerProThrValGlnLeuGln ProLeuLeuProAspProSerAlaIleThrProAsnSerThrSerProLeuLeuPheAlaAlaHisProHisPheGlnAsnLeuSerGlnGluGlu 320 280 310 240 801

501 ATTCCACACACTGAACAAGCTCCCAAACAGGACCTCTGGATTGTTGGAGACAAACTCCAGTATCTCAGCCAGAACTACTGGCTCTGGATTTCTCAAGAGG PheHisThrLeuAsnLysLeuProAsnArgThrSerGlyLeuLeuGluThrAsnSerSerIleSerAlaArgThrThrGlySerGlyPheLeuLysArg

1001 GTGCTCAGACCCTGCCAACTTCAGCA

30/85 SUBSTITUTE SHEET (RULE 26)

Ser Pro Ala Pro Pro Ala Cys Asp Pro Arg Leu Leu Asn Lys Leu Leu Arg Asp Ser His Val Leu His Gly Arg Leu Ser Gln Cys Pro Asp Ile Asn Pro 

9

LeuSerThrProValLeuLeuProAlaValAspPheThrLeuGlyGluTrpLysThrGlnThrGluGlnThrLysAlaGlnAspValLeuGlyAlaThr CTTTGTCCACACCTGTCCTGCTGCTGTGGACTTCACCTTGGGAGAATGGAAAACCCAGACGGAGCAGACAAAGGCACAGGATGTCCTGGGAGCCAC 40 101

20

LeuGlyAlaLeuGlnAspLeuLeuGlyMetGlnGlyArgThrThrAlaHisLysAspProSerAlaIlePheLeuAsnPheGlnGlnLeuLeuArgGlyLys AACCCTTCTGCTGGAGGCAGTGATGACAGCACGGGGACAAGTGGGACCCCCTTGCCTCTCATCCCTGCTGGTGCAGCTTTCTGGACAGGTTCGCCTCCTC ThrLeuLeuLeuGluAlaValMetThrAlaArgGlyGlnValGlyProProCysLeuSerSerLeuLeuValGlnLeuSerGlyGlnValArgLeuLeu 110 201

ValArgPheLeuLeuLeuValValGlyProSerLeuCysAlaLysArgAlaProProAlaIleAlaValProSerSerThrSerProPheHisThrLeu 160 140

401 AGGTGCGTTTCCTGCTTCTTTGTAGTGGGCCCTCCCTCTGTGCCAAGAGGGCCCCACCCGCCATAGCTGTCCCGAGCAGCACCTCTCCATTCCACACACT

FIG.2 1 A

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**AsnLysLeuProAsnArgThrSerGlyLeuLeuGluThrAsnSerSerIleSerAlaArgThrThrGlySerGlyPheLeuLysArgLeuGlnAlaPhe** 501 GAACAĀGCTCCCAAACAGGACCTCTGGĀTTGTTGGAGACAAACTCCAGTATCTCAGCCAGĀACTACTGGČTCTGGĀTTTCTCAĀGAGĞCTGCAGGCATTC 230

ArgAlaLysIleProGlyLeuLeuAsnGlnThrSerArgSerLeuAspGlnIleProGlyHisGlnAsnGlyThrHisGlyProLeuSerGlyIleHisGly 601 AGÁGCCAAGATTCCTGGTCTGAACCAAACCTCCAGGTCCCTAGACCAAATCCCTGGACACCAGAATGGGACACACAGGACCTTGAGTGGAATTCATG

LeuPheProGlyProGlnProGlyAlaLeuGlyAlaProAspIleProProAlaThrSerGlyMetGlySerArgProThrTyrLeuGlnProGlyGlu 701 GACTCTTTCCTGGÁCCCCAACCCGGGGCCCTCGGAGCTCCAGACATTCCTCCAGCAACTTCAGGCATGGGCTCCCGGCCAACCTÁCCTCCAGCCTGGÁGA

SerproSerProAlaHisProSerProGlyArgTyrThrLeuPheSerProSerProThrSerProSerProThrValGlnLeuGlnProLeuLeuPro

290

GACCCTCTGCGATCACACCCAACTCTACCAGTCCTCTTCTATTTGCAGCTCACCCTCATTTCCAGAACCTGTCTCAGGAAGAAGTAAGGTGCTCAGAACCC AspproSerAlaIleThrProAsnSerThrSerProLeuLeuPheAlaAlaHisProHisPheGlnAsnLeuSerGlnGluGlu 320 801

1001 TGCCAACTTCAGCA

901

32/85 SUBSTITUTE SHEET (RULE 26)

pML pML2	1 SPAPPACDPRILNKLLRDSHVLHGRLSQCPDINPLSTPVLLPAVDFT 2 1 SPAPPACDPRILNKLLRDSHVLHGRLSQCPDINPLSTPVLLPAVDFT	L G E
pML pML2	51 WKTQTEQTKAQDVLGATTLLLEAVMTARGQVGPPCLSSLLVQLSGQV 2 51 WKTQTEQTKAQDVLGATTLLLEAVMTARGQVGPPCLSSLLVQLSGQV	RLL
pML pML2	101 LGALQDLLGMQLPPQGRTTAHKDPSAIFLNFQQLLRGKVRFLLLVVG 2 101 LGALQDLLGMQGRTTAHKDPSAIFLNFQQLLRGKVRFLLLVVG	P S L
pML pML2	151 CAKRAPPAIAVPSSTSPFHTLNKLPNRTSGLLETNSSISARTTGSGF 2 147 CAKRAPPAIAVPSSTSPFHTLNKLPNRTSGLLETNSSISARTTGSGF	L K R
pML pML2	201 L QAFRAK IPGLL NOTSRSL DQIPGHQNGTHGPLSGIHGLFPGPQPGA 2 197 L QAFRAK IPGLL NOTSRSL DQIPGHQNGTHGPLSGIHGLFPGPQPGA	1 L G A
pML pML2	251 PDIPPATSGMGSRPTYLQPGESPSPAHPSPGRYTLFSPSPTSPSPTV 2 247 PDIPPATSGMGSRPTYLQPGESPSPAHPSPGRYTLFSPSPTV	0 1 0
pML pML2	PLLPDPSAITPNSTSPLLFAAHPHFONLSOEE FIG. 22 297 PLLPDPSAITPNSTSPLLFAAHPHFONLSOEE	

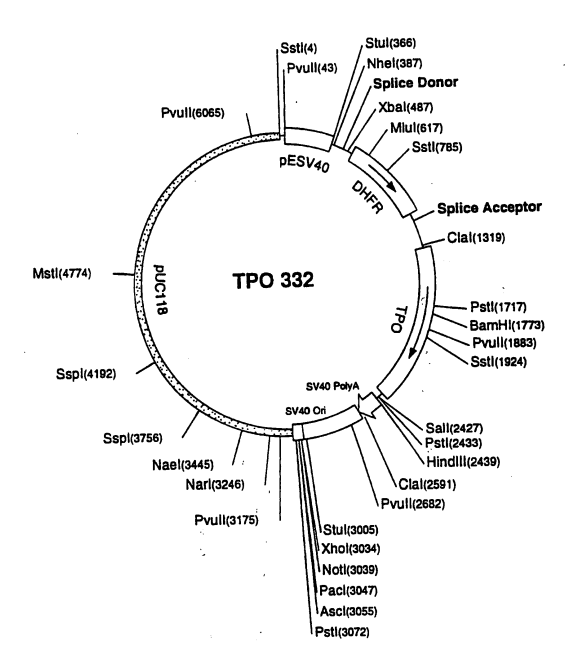
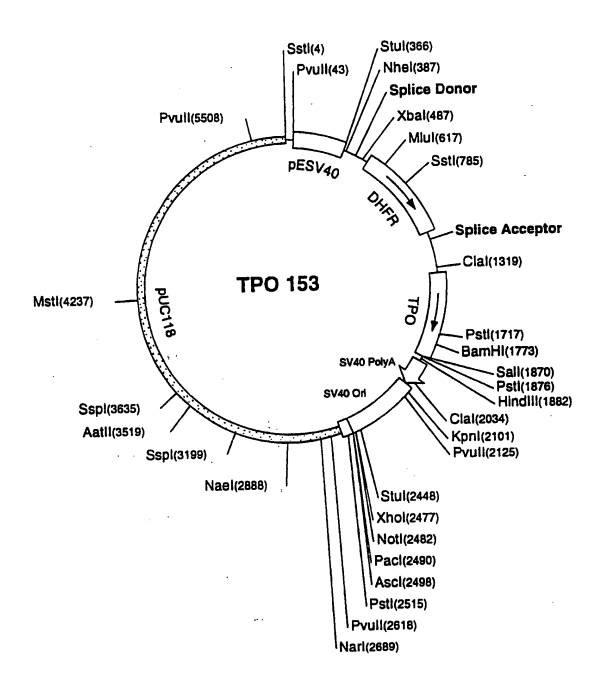


FIG.23

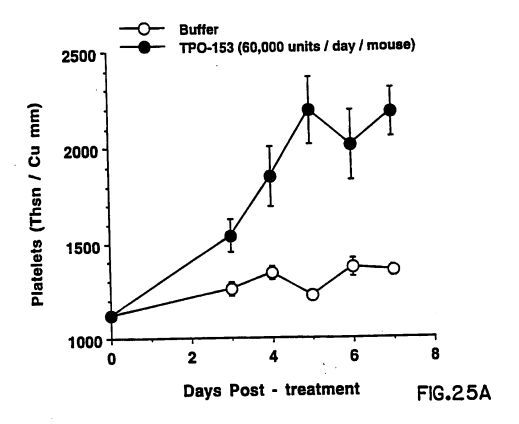
34/85

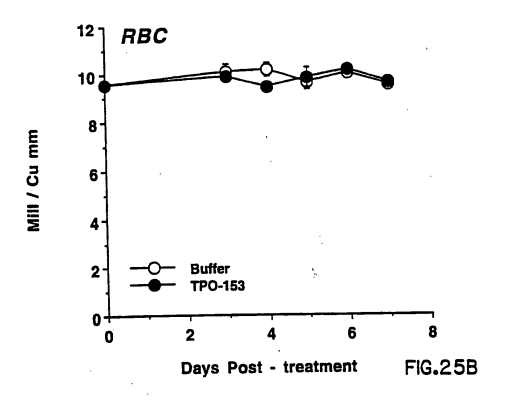


**FIG.24** 

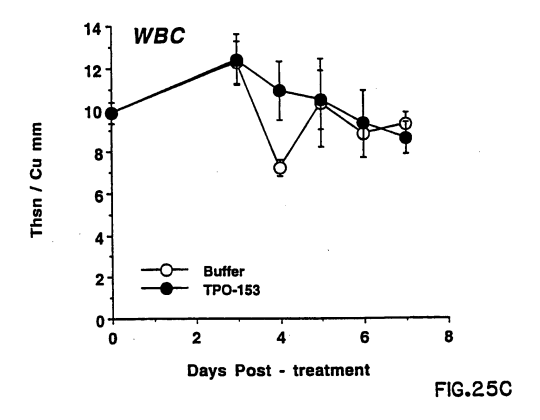
35/85

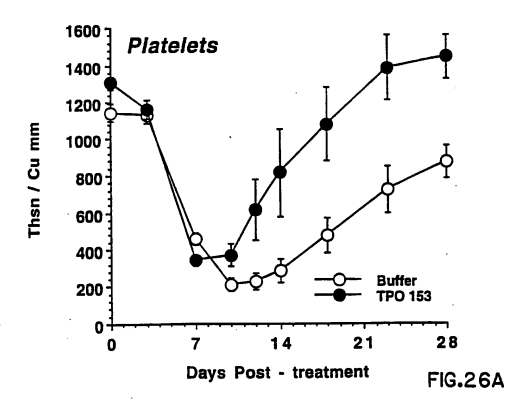
PCT/US94/14553





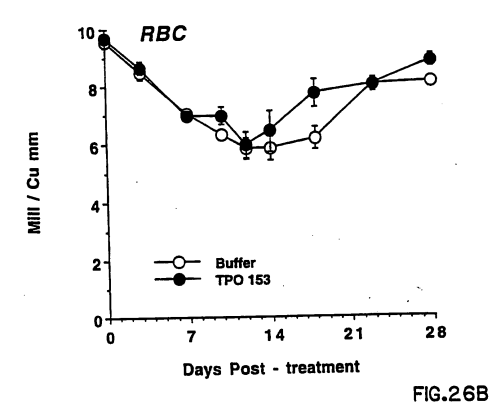
36/85 SUBSTITUTE SHEET (RULE 26)

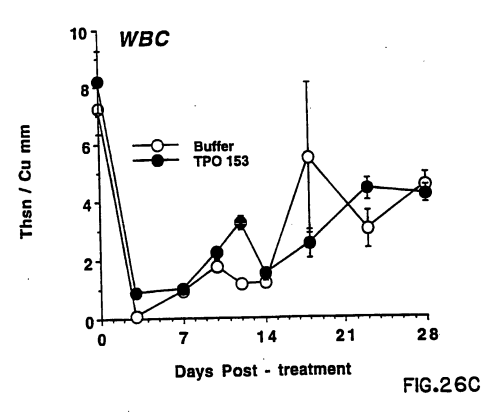




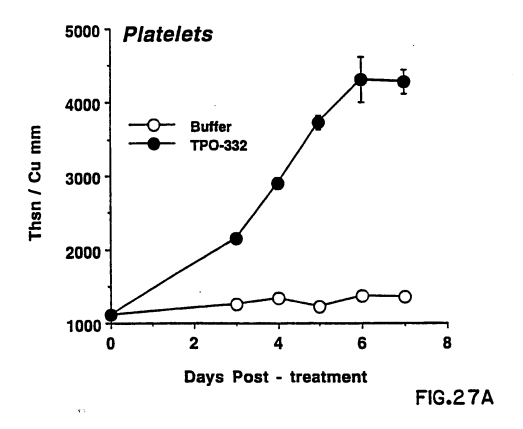
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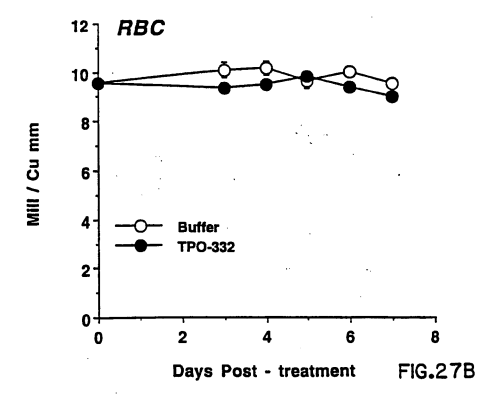
SUBSTITUTE SHEET (RULE 26)



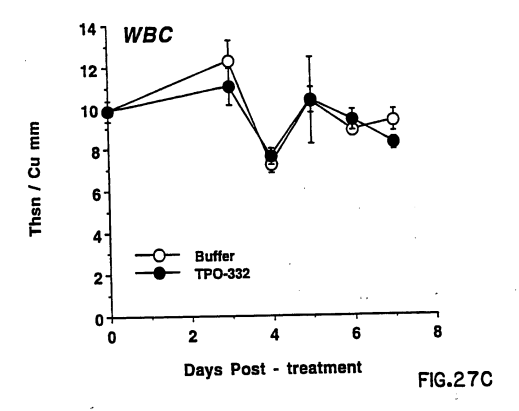


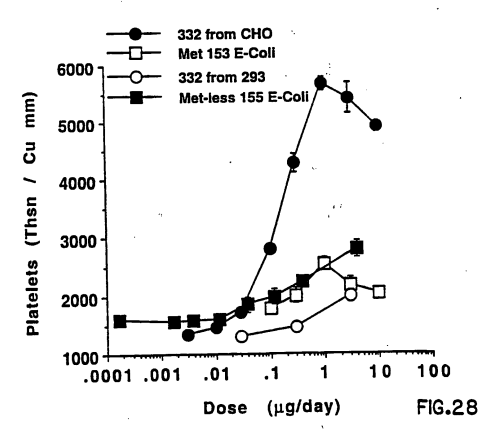
38/85
SUBSTITUTE SHEET (RULE 26)





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SUBSTITUTE SHEET (RULE 26)





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SUBSTITUTE SHEET (RULE 26)

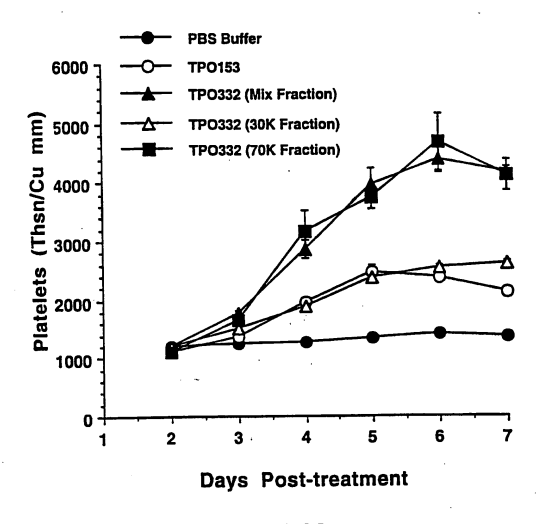
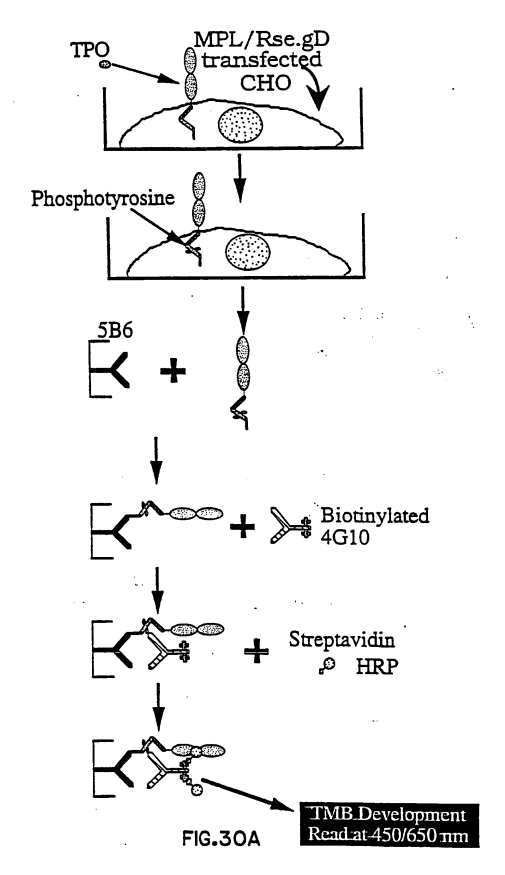


FIG.29

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PCT/US94/14553



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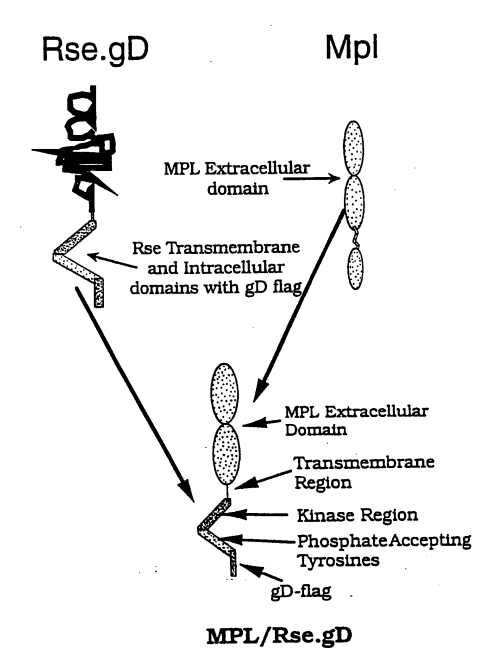


FIG.30B

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FIG.31

5x10<sup>4</sup>MPL/Rse.gD cells per well in 96-well culture plates (flat-bottom) Stimulate w/ ligand x 30 minutes Stimulation may be done in either 50 or 100 µl Solublize in: 150 mM NaCl 0.5 % Triton-X 100 50 mM HEPES, pH 7.5 Na<sub>3</sub>VO<sub>4</sub> (phosphatase inhibitor) Leupeptin, Aprotinin & AEBSF (protease inhibitors) Transfer 85 µl of cell lysate to ELISA well coated with capture antibody, 5B6 (anti-gD peptide mAb) incubate on shaker x 2 hr Wash plate and add biotinylated-4G10. 400 pg/ml, 100 μl/well. Incubate on shaker x 2 hr Wash plate and add Zymed streptavidin/HRP 1:10K, 100 µl/well. Incubate on shaker x 30 min Wash plate and add Kirkegaard & Perry TMB substrate, 100 µl/well. Develop x 10 min

Stop with 1 M H<sub>3</sub>PO<sub>4</sub> and read at 450/650 nM

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SUBSTITUTE SHEET (RULE 26)

			H						
	CAGTTAGGGT GTCAATCCCA	SCLF1	ecoRII	dsaV	DSCNI	apyı	sexAI	CTCAATTAGT CAGCAACCAG	Greerregre
	GAATGTGTGT CAGTTAGGGT CTTACACACA GTCAATCCCA		ralli						TICGIACGIA GAGIIAAICA GICGIIGGIC
aluI sau3AI pvuII mboI/ndeII[dam-] dpnI[dam+] vuI/bspCI dpnII[dam-] taqI[dam-] icrI nspBII	GACAGCTGTG	staNI ppu10I	nsil/avallI	nlaIII	sphI	Idsu	IHdsu	AAGCATGCAT	TTCGTACGTA
aluI sau3AI pvuII mboI/ndeII[cd dpnI[dam+] pvuI/bspCI pleI dpnII[dam-] hinfI taqI[dam-]	GAG							CCCAGGCTCC CCAGCAGGCA GAAGTATGCA AAGCATGCAT	CTTCATACGT
rmaI	ATTATTGACT TAATAACTGA	,						CCAGCAGGCA	GGTCGTCCGT
pHI	CCCGACATTG GGGCTGTAAC nlaIV	SCLFI	PCORTI	dsaV	bstNI	apvI[dcm+]	bsaJI	CCCAGGCTCC	GGGTCCGAGG
ssti saci hgiJII hgiAI/aspHI ecl136II bsp1286 bsiHKAI bmyI	taq1 TTCGAGCTCG AAGCTCGAGC							GTGGAAAGTC	CACCTTTCAG
	-							71	4

FIG.32

FIG.32B

```
nlaIII
                                                                                                                                                                                                                                                                                                                                                  GCTGACTAAT TTTTTTTATT TATGCAGAGG CCGAGGCCGC CTCGGCCTCT GAGCTATTCC AGAAGTAGTG
                                                                                                                                                                                                                                                                                                                                                              CGACTGATTA AAAAAATAA ATACGTCTCC GGCTCCGGCG GAGCCGGAGA CTCGATAAGG TCTTCATCAC
                                                                                                                                                                                                                       TATCAGGGCG GGGATTGAGG CGGGTAGGGC GGGGATTGAG GCGGGTCAAG GCGGGTAAGA GGCGGGGTAC
                                                                                                                                                                                                          CCGCCCCATG
                                                                                              GTCAGCAACC
                                                                                                          CACACCTTTC AGGGGTCCGA GGGGTCGTCC GTCTTCATAC GTTTCGTACG TAGAGTTAAT CAGTCGTTGG
                                                                                                                                                                                            acil bsaJI
                                                                                                                                                                 ncol
                                                                                                                                                     styI
                                                                                                                                                                                bslI dsaI
                                                                                                                                                                                                            211 ATAGICCCGC CCCIAACICC GCCCAICCCG CCCCIAACIC CGCCCAGIIC GCCCAIICI
                                                                                                GIGIGGAAAG ICCCCAGGCI CCCCAGCAGG CAGAAGIAIG CAAAGCAIGC AICICAAIIA
                                                                                                                                                                                                                                                                                                                           aluI
                                                                                                                                                                                                                                                                                                                                         haeIII/palI
                                                                                                                                                                                                 acil
                                                                                                                                                                                                                                                                                                              ddeI
                 ppu10I
                                                                                                                                                                                                                                                                                                                          mnlI
                                                                                     IHdsu
                                                                        nspl
                                                            sphI
                                                                                                                                                                                                   acil bsrI
                                 nsil/avallI
                                                                                                                                                                                                                                                                                                  haeIII/palI
                                                                                                                                                                                                                                                                                                                             haeIII/palI bsaJI
                                                                                                                                                                                                                                                                                                                mnlI
        sfaNI
                                                                                                                                                                                                                                                                                                                                            mnlI bsaJI aciI
                                                                                                                                                                                                                                                          fnu4HI
                                               nlaIII
                                                                                                                                                                                                                                                                        bglI
                                                                                                                                                                                                                                                                                      sfiI
                                                                                                                                                                                                                                                                                                                 mnlI
                                                                                                                                                                                         acil
                                                                                                                                                                                                       acil fokl
                                                                                 apyI[dcm+]
nlaIV
                                                                  bstNI
                                                                                             bsaJI
                           mvaI
                                         ecoRII
              SCLFI
                                                       dsaV
                                                                                    [dcm+]
                                                                                                                141
```

GTACCGCCTA TAGAGCGATA CATGGCGGAT ATCTCGCTAT eagI/xmaIII/eclXI scfI haeIII/palI acil csp6I donor rsal mcrI eaeI cfrI hpaII GCGCCTAAGG GGCACGGTTC TCACTGCATT Idsm AGTGACGTAA GCTTATCCGG ^splice CGAATAGGCC maeII maeIII aluI maeI rmaI TTTGGAGGCC TAGGCTTTTG CAAAAAGCTA CCGTGCCAAG TCCTCCGAAA AAACCTCCGG ATCCGAAAAC GTTTTCGAT nheI aluI fnuDII/mvnI CGCGGATTCC hinFI tfiI **bsh1236I** haeIII/palI aciI bstul thaI rmaI mnl1 maeI bsaJI avrII styl blnI CCGGGAACGG TGCATTGGAA GGCCCTTGCC ACGTAACCTT haeI stuI AGGAGGCTTT mnlI SCrFI hpall caulI dsaV Idsm nciI 351 401

DHFR ATG' FIG.32C

CCAAAATATG

pflMI bslI GGTTTTATAC

TTGACGTAGC AGCGGCACAG

TAGGGGCGAC GGTAGTACCA AGCTGGTAAC

AGAGGATTIT ATCCCGCTG CCATCATGGT

TCTCCTAAAA

acil

TCGACCATIG AACTGCATCG TCGCCGTGTC

sfaNI

taqI

nlaIII

fnu4HI

bbvI nspBII

mnll

bslI

```
ddeI
                                                                                                                                                                                                                        GGTGTTGGAG AAGTCACCTT CCATTTGTCT TAGACCACTA ATACCCATCC TTTTGGACCA AGAGGTAAGG
                                                                                                                                                                                                              CCACAACCIC TICAGIGGAA GGIAAACAGA AICIGGIGAI IAIGGGIAGG AAAACCIGGI ICICCAIICC
                                                                                             GGAACGAGIT CAAGIACIIC CAAAGAAIGA
                                                                                                        CCTTGCTCAA GTTCATGAAG GTTTCTTACT
                                                                                                                                                                                         apyI[dcm+]
                                                                                                                                                       ecoRII
                                                                                                                                                                             bstNI
                                                                                                                                 scrFI
                                                                                                                                                                   dsaV
                                                                                                                                            mvaI
                                                                                                                                                                                                     sexAI
                                                                          csp6I
                                                               rsal
                                                                                      scal
                                                                                     asp700
                                                                            XmnI
                                                                                        ddeI
                                                                                                              CCCTAACCGT TCTTGCCTCT GGATGGGACC GGAGGCGAGT
                                                                                                   CCTACCCTGG CCTCCGCTCA
                                                                                                                                                                                                        hphI
                                bsrBI
haeIII/palI
                                                                  acil
                                                                                         mnlI
                                                                              apyI[dcm+]
                                                                                                                                                                                                hinfI
                                                                                                                                                                                    tfiI
                                                                                                                                                                                                          alwNI
            haeI
                                             ecoRII
                                                                    bstNI
                       scrFI
                                                                                         bsaJI
                                   mvaI
                                                          dsaV
                                                                                 bsmAI
                                                                                            bsaI
                                                                                                       GGGATTGGCA AGAACGGAGA
                                                                                                                                                                                                 earI/ksp632I
                                                                                                                                                                           eco57I
                                                                                                                                                                                       Ilodm
                                                                                                            541
                                                                                                                                                                                                                           611
```

ACTCTTCTTA GCTGGAAATT TCCTGTCTTA ATTATATCAA GAGTCATCTC TTGAGTTTCT TGGTGGTGCT FIG.32D

TGAGAAGAAT ČGACCTTTAA AGGACAGAAT TAATATAGTĪ CTCAGTAGAG AACTCAAAGA ACCACCACGA

aseI/asnI/vspI

ahaIII/draI

tru9I mseI

tru9I

mseI

hinfI tfiI

TTGGCAAGTA	haeIII/palI haeI		mvaI		
sstI sac! hgiJII hgiAI/aspHI ecl136II bsp1286 bsiHKAI bmyI banII bstXI fokI sfaNI msel bsaWI aluI bstXI fokI sfaNI msel bsaWI 751 GGAGCTCATT TTCTTGCCAA AAGTTTGGAT GATGCCTTAA GACTTATTGA ACACCGGAA TTGGCAAGTA CCTCGAGTAA AAGAACGTT TTCAAACCTA CTACGGAATT CTGAATAACT TGTTGGCCTT AACCGTTCAT		scrFI	TIII	ecoRII	
I /bfrI GACTTATTGA CTGAATAACT		scrFI	mvaI	ecoRII	
tru9I af111/bfrI fokI sfaNI mseI GGAT GATGCCTTAA GACT CCTA CTACGAATT CTGA					
fok I PAGTTTGGAT TTCAAACCTA					
I bstXI TTCTTGCCAA AAGAACGGTT					
sstl saci hgiJII hgiAI/aspHI ecl136II bsp1286 bsiHKAI bmyI banII aluI GGAGCTCATT T					'.
751					

apyl[dcm+] hinfl apyl[dcm+] cCAGGAAGCC ATGAATCAAC CAGGCCACCT GGTCCTTCGG TACTTAGTTG GTCCGGTGGA apyI[dcm+] hinfI nlaIII bstNI TTCATCTGTA CCAAACCTAT CAGCCTCCGT CAAGACAAAT GTTCTGTTTA 821 AAGTAGACAT GGTTTGGATA GTCGGAGGCA mnlI accI nlaIII

bstNI

dsaV

tfil

dsaV

GTGACAAGGA TCATGCAGGA ATTTGAAAGT GACACGTTTT TCCCAGAAAT TGATTTGGGG ATCTGAGAAA CACTGTTCCT AGTACGTCCT TAAACTTTCA CTGTGCAAAA AGGGTCTTTA ACTAAACCCC maeII aflIII maeIII apoI mboI/ndeII[dam-] dpnII[dam-] nlaIII dpnI[dam+] maeIII alwI[dam-] sau3AI TAGACTCTTT hinfI pleI 891

hgal
hinll/acyl
ahall/bsaHl
scrfl
mval mnll
ecoRII
dsaV
bstNI ecoNI

apyl[dcm+] mn bsaJI bslI ddeI

bsaji dde. 961 AAATATAAAC CTCTCCCAGA ATACCCAGGC GTCCTCTGTG TTTATATTTG GAGAGGGTCT TATGGGTCCG CAGGAGAGAC

FIG.32F

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1001 AGGTCCAGGA GGAAAAAGGC ATCAAGTATA AGTTTGAAGT CTACGAGAAG AAAGACTAAC AGGAAGATGC TCCAGGICCT CCTTTTTCCG TAGTTCATAT TCAAACTTCA GATGCTCTTC TTTCTGATTG sfaNI apyI[dcm+] mnlI ecoRII bstNI dsaV mvaI sau96I avall asuI

SCLFI

nlaIII styl

TCCTTCTACG

· END DHFR

Ilodm

Ilodm

accI

sfaNI

ncol

dsaI

bsaJI nsil/avaIII ppu10I aluI

mnlI

1071 TITCAAGTIC ICTGCICCC ICCTAAAGCI AIGCAITITI AIAAGACCAI GGGACTITIG AAAGIICAAG AGACGAGGGG AGGAITICGA IACGIAAAAA IAITCIGGIA CCCIGAAAAC

CAGGIGICCA CICCCAGGIC GTCCACAGGT GAGGGTCCAG

1201 ATACGATTTA GGTGACACTA TAGATAACAT CCACTTTGCC TTTCTCTCCA

scfI

hphI

TATGCTAAAT CCACTGTGAT ATCTATTGTA GGTGAAACGG AAAGAGAGT

styl

apyI[dcm+] sau96I avalI asuI TGCGCCGATG TTAATTATGT ATTGGAATAC ATAGTATGTG ACGCGGCTAC AATTAATACA TAACCTTATG TATCATACAC ecoRII bstNI SCrFI dsaV mvaI bsaJI bslI aseI/asnI/vspI fnuDII/mvnI tru9I mseI bsh1236I fnu4HI bstUI aciI thaI fokI GACCGAAATC TAGGGGAACC GAAGCAATCT CTTCGTTAGA mboI/ndeII[dam-] CTGGCTTTAG ATCCCCTTGG bsaJI dpnII[dam-] bstYI/xhoII dpnI[dam+] alwI[dam-] maeIII sau3AI 1131

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mbo1/ndeI1[dam-] xmaI/pspAI scrFI hpall dpnII[damdpnI[dam+] nlaIV cauII bamHI bsaJI Idsm bstYI/xhoI] nciI dsaV hincII/hindII alwI[damscrFI caulI bsaJI smaI dsaV avaI alwI[dam-] nciI acci xbai mnli bsaJi GICGACICIA GAGGAICCCC CAGCTGAGAT CTCCTAGGGG sau3AI taqI rmaI salI maeI hinfI pleI DSPMI 1271 CAACTGCACC TCGGTTCTAA GCTTCTGCAG GTTGACGTGG AGCCAAGATT CGAAGACGTC pstI hindIII aluI scfI ddeI bsaJI mnlI

FIG.321

TATCGTAGTG

TITAAAGTGT TTATTTCGTA AAAAAGTGA CGTAAGATCA ACACCAAACA GGTTTGAGTA TGTGGTTTGT CCAAACTCAT ATCGATGGCC GCCATGGCCC AACTTGTTTA TTGCAGCTTA TAATGGTTAC AAATAAAGCA TIGAACAAAT AACGICGAAT ATTACCAAIG TTTATTTCGT maeIII GCATTCTAGT rmaI bsmI maeI aluI fnu4HI ^sv40 early poly A bbvI AAATITCACA AATAAAGCAI ITITITCACI haeIII/palI sau96I TAGCTACCGG CGGTACCGGG asuI nlaIII taqI haeIII/palI dsaI styl ncol fnu4HI acil bglI sfil cfrI eaeI apoI 1391 ATAGCATCAC CCCCTTAAGT GGGGAATTCA sfaNI ecoRI apoI 1321

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FIG.32K

```
CGGCGCAGCA CCATGGCCTG AAATAACCTC TGAAAGAGGA ACTTGGTTAG GTACCTTCTG AGGCGGAAAG
                                                                                                                                                                                                                                                                                                                                          GCCGCGTCGT GGTACCGGAC TTTATTGGAG ACTTTCTCCT TGAACCAATC CATGGAAGAC TCCGCCTTTC
                                                                                                                                                                                                                                                                                                                 aciI
                                                                                                                                                                                                                                                                                                     mnlI
                                                                                                                                                                                                                                                                                                                 ddeI
                                                                                                                                                                                                                                     csp6I
                                                                                                                                                                                                                         rsaI
                                                                                                                                                                                                                                                                                                   asp718
                                                                                                                                                                                                                                                                                                                 acc651
                                                                                                                                                                                                                                                                          hgiCI
                                                                                                                                                                                                                                                nlaIV
                                                                                                                                                                                                                                                              kpnI
                                                                                                                                                                                                                                                                                        banI
                                                                                                                                            aseI/asnI/vspI
                                                                                                                                                                     GGATCGATCG GGAATTAATT
                                                                                                                                                                                 CCTAGCTAGC CCTTAATTAA
             mboI/ndeII[dam-]
                                                                                                                                                                                                                                                                                                                   mnlI
                                                                           tru9I
                                                                                                       mseI
                                                                                        claI/bsp106[dam-]
                                                                                                                                                         asp700
                                                                                                                  mboI/ndeII[dam-]
                                                                                                                                dpnI[dam+] xmnI
                                     dpnII[dam-]
                       dpnI[dam+]
                                                  pvuI/bspCI
                                                                           taqI[dam-]
                                                                                                                                                                                                 sv40 origin^
                                                                                                                                            dpnII[dam-]
sau3AI
                                                                                                                                                         alwI[dam-]
                                                                                                                                                                                                                                                                                                                   mnll
                                                              mcrI
                                                                                                    sau3AI
                                                                                                                                                                                                                                       haeIII/palI
                                                                                                                                                                                    GTTACATAGA ATAGTACAGA
                                                                                                                                                                      CAATGTATCT TATCATGTCT
                                                                                                                                                          nlalII
                                                                                                                                                                                                                                                   haeI
                                                                                                                                                                                                                                                                                                                    hhal/cfoI nlaIII
                                                                                                                                                                                                                                                                                                       bsaJI
                                                                                                                                                                                                                                                                 styI
                                                                                                                                                                                                                                                                             ncol
                                                                                                                                                                                                                                                                                          dsaI
                                                                                                                                                                                                                                                                             fnu4HI
                                                                                                                                                                                                                                                                                          bbvI
                                                                                                                                                                                                                                                                                                       hinPI
                                                                                                                                                                        1461
                                                                                                                                                                                                                                                                                                                                  1501
```

nlaIV scrFI mvaI ecoRII	bstni apyl[dcm+] bsaJI ccccAGGCTC CCCAGCAGGC AGAAGTATGC GGGGTCCGAG GGGTCGTCCG TCTTCATACG	mval bstNI apyI[dcm+] bsaJI GTCCCCAGGC GCAGAAGTAT CAGGGGTCG AGGGGTCGTC CGTCTTCATA	acil caragreece ceceraacre egecearer grareagge gegearreag ecegeragg
	TGTGGAAAGT ACACCTTTCA	scrFI mval ecoRII dsaV bstNI apyI[dcm+] exAI cca GGTGTGGAAA	aciI CATAGTCCCG (
	TCAGTTAGGG TGTGGAAAGT AĜTCAATCCC ACACCTTTCA	scrFI mval ecoRII dsaV bstNI apyI[dcm+] sexAI TCAGCAACCA GGTGTGGAAA AGTCGTTGGT CCACCCTTT	aNI 01 /avaIII I CATCTCAATT AGTCAGCAAC GTAGAGTTAA TCAGTCGTTG
FIG.32L	GGAATGTGTG CCTTACACAC	TCTCAATTAG	W
	aluI pvuII nspBII 1571 AACCAGCIGT TTGGTCGACA	sfaNI ppu10I nsi1/avaIII nlaIII sphI nspII nspHI TTTCGTACGT	sfaN ppu10I nsiI/a nlaIII sphI nspI nspHI nspHI CGTTTCGTAC GT

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mnlI CCGCCCAGIT CCGCCCAIIC ICCGCCCCAI GGCIGACIAA IITITITIAI ITAIGCAGAG GGCGGGTCAA GGCGGGTAAG AGGCGGGGTA CCGACTGATT AAAAAAATA AATACGTCTC nlaIII acil bsaJI styI ncol bslI dsaI acil bsrI asiI

1781

haeIII/palI avrII blnI stuI haeIII/palI fnu4HI bglI sfiI

bsaJI

hael

mnlI

aluI

bsaJI mnlI

haeIII/palI

ddeI

mnlI

mnlI

styI

CGGCTCCGGC GGAGCCGGAG ACTCGATAAG GTCTTCATCA CTCCTCCGAA AAAACCTCCG GAGGAGGCTT TTTGGAGGC mnlI mnlI GCCGAGGCCG CCTCGGCCTC TGAGCTATTC CAGAAGTAGT haeIII/palI bsaJI aciI 1841

FIG.32M

tru9I GCACTGGCCG TCGTTTACA ACGTCGTGAC TGGGAAAACC CTGGCGTTAC CCAACTTAAT CGTGACCGGC AGCAAATGT TGCAGCACTG ACCCTTTTGG GACCGCAATG GGTTGAATTA GTIACCTCGA GCGGCCGCTI AATTAAGGCG CGCCATTIAA ATCCTGCAGG GATCCGAAAA CGTTTTTCGA CAATGGAGCT CGCCGGCGAA TTAATTCCGC GCGGTAAATT TAGGACGTCC sse8387I pspMI scfI bsdI pstI ahaIII/draI ^linearization linker inserted into Hpal site maeIII hhal/cfol tru91 msel tru91 bsh1236I msel msel bssHII swaI apyI[dcm+] fuuDII/mvnI hhaI/cfoI ecoRII bstNI scrFI bsaJI dsaV mvaI hinPI bstuI thaI hinPI ascI eag1/xmaIII/eclXI tru9I bsrľ haeIII/palI pacI maeIII aluI maeIII bsrBI fnu4HI aciI maeII fnu4HI eaeI paeR7I cfrI mcrI aciI notI taqI avaI xhoI mnlI ^start pUC118 haeIII/palI eaeI cfrI 1971 TAACAGCTTG ATTGTCGAAC aluI maeIII rmaI maeI 1901

mbol/ndeII{dam-] dpnII[dam-] pvuI/bspCI dpnI[dam+] mcrI CGCCTTGCAG CACATCCCCC CTTCGCCAGC TGGCGTAATA GCGAAGAGGC CCGCACCGAT GCGGAACGIC GIGIAGGGGG GAAGCGGICG ACCGCAITAI CGCIICICCG GGCGIGGCIA sau3AI acil earI/ksp6321 sau96I haeIII/palI mnlI Iloqm nspBII aluI IInvd fokI fnu4HI bbvI

hhal/cfol nlaIV narI kasI hinlI/acyI hgiCI haeII aciI banI sfaNI

hinPI

2101 CGCCCTTCCC AACAGTTGCG TAGCCTGAAT GGCGAATGGC GCCTGATGCG GTATTTTCTC CTTACGCATC GAATGCGTAG GCGGGAAGGG TTGTCAACGC ATCGGACTTA CCGCTTACCG CGGACTACGC CATAAAAGAG bglI

FIG.320

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hhaI/cfoI hinPI fnuDII/mvnI bstUI scfI bsh1236I hinPI thaI

fnu4HI rsal hhal/cfol

TGTGCGGTAT TTCACACCGC ATACGTCAAA GCAACCATAG TACGCGCCCT GTAGCGGCGC ACACGCCATA AAGTGTGGCG TATGCAGTTT CGTTGGTATC ATGCGCGGGA CATCGCCGCG aciI bslI csp6I maeII

acil

acil

2171

hhaI/cfoI fnu4HI hinPI fnuDII/mvnI fnu4HI thal

fnuDII/mvnI thaI

bsh1236I bstUI

aciI

bsrBI

hhaI/cfoI haell mael

haeII

hinPI

rmaI

acil

hhaI/cfoI

hinPI

maeIII bbvI

maeIII

mseI bsh1236I

TAATTCGCGC

2231

tru9I aciI

TEGITACECE CAGCGIGACC GCIACACITE CGCCCACACC ACCAATGCGC GTCGCACTGG CGATGTGAAC ATTAAGCGCG GCGGGTGTGG

GGTCGCGGA TCGCGGGCGA

CCAGCGCCT AGCGCCCGCT

bsp1286 banII bmyI aluI hpaII Idsm

hgiJII nlaIV

> maeII cfr10I naeI

GGAAAGCGAA AGAAGGGAAG GAAAGAGCGG TGCAAGCGGC CGAAAGGGGC AGTTCGAGAT TTAGCCCCCG ACGITCGCCG GCTITCCCCG TCAAGCICIA AAICGGGGGC TCTTCCCTTC CTTTCTCGCC mbolI CCTTTCGCTT 2301

FIG.32P

60/85

hinPI acil

bctUI

hhaI/cfoI

Ilum !!

nlaIV

hgiCI

tadI banI

nlaIV

2371

hphI

CTTGATTTGG GGGGTTTTTT GAACTAAACC CCCCAAAAA CCGTGGAGCT GGCACCTCGA AGGGAAATCC CAAGGCTAAA TCACGAAATG TCCCTTTAGG GTTCCGATTT AGTGCTTTAC

.haeIII/palI maeII

sau96I asuI

drallI bsaAI

2401

:

maeII pleI

hinfI drdI

bslI

bslI

GIGAIGGIIC ACGIAGIGGG CCAICGCCCI GAIAGACGGI IIITCGCCCI IIGACGIIGG AGICCACGII CACTACCAAG TGCATCACCC GGTAGCGGGA CTATCTGCCA AAAAGCGGGA AACTGCAACC TCAGGTGCAA

bsrI hinfI pleI

GGACTCTTGT TCCAAACTGG AACAACACTC AACCCTATCT CGGGCTATTC TTTTGATTTA CCTGAGAACA AGGTTTGACC TTGTTGTGAG TTGGGATAGA GCCCGATAAG AAAACTAAAT avaI GAAATTATCA CTTTAATAGT

tru9I tru9I

tru9I

mseI

AGCTGATTTA ACAAAATTT apoI mseI aluI TTAAAAATG mseI haeIII/palI GGCCTATTGG TAAGGGATTT TGCCGATTTC

AATTTTTAC TCGACTAAAT TGTTTTTAAA CCGGATAACC ATTCCCTAAA ACGGCTAAAG

2501

tru9I mseI

thal fuuDII/mvnI maeII bsp1286 fuuDII/mvnI maeII bsiHKAI apoI tru9I tru9I apaLI/snoI rsaI bsh1236I mseI sspI mseI AACGCGAATT TTAACAAAAT ATTAACGTTT ACAATTTTAT GGTGCACTCT CAGTACAATC TTGCGCTTAA AATTGTTTTA TAATTGCAAA TGTTAAAATA CCACGTGAGA GTCATGTTAG	bsrI hinPI maeIII fnu4HI NI acil mseI acil bsaAI tth1111/aspI bbvI G CCGCATAGTT AAGCCAACTC CGCTATCGCT ACGTGACTGG GTCATGGCTG CGGGGGCTGT C GGCGTATCAA TTCGGTTGAG GCGATAGCGA TGCACTGACC CAGTACCGAC GCGGGGCTGT	hinPI hhal/cfol thal thal thal fluDII/mvnI bstUI  acil acil hgal drdI acil acil hgal drdI acil GGCGTTGTCTG CTCCCGGCAT GCGTTACAG ACAAGCTGTG GGGCGTTGT GCGCACTGC CCGAACAGAC GAGGGCCGTA GGCGAATGTC TGTTCGACAC GGGCGGTTGT GGGCGACTGC CCGAACAGAC GAGGGCCGTA GGCGAATGTC TGTTCGACAC
thal fnuDII/mvnI apol bstUI tr bsh1236I ms AACGCGAATT TT	fnu4HI sfaNI aciI TGCTCTGATG CCGC ACGAGACTAC GGCG	aciI CCCGCCAACA GGGCGGTTGT
2631	2691	2761

FIG

hhaI/cfoI

CTTTACACGC

2951 ATAATAATGG TTTCTTAGAC GTCAGGTGGC ACTTTTCGGG GAAATGTGCG

aatii ddei maeii TATTATTACC AAAGAATCTG CAGTCCACCG TGAAAAGCCC

hinPI

TGGCAGAGGC CCTCGACGTA CACAGTCTCC AAAAGTGGCA GTAGTGGCTT TGCGCGCTCC GTCATAAGAA CAGTATTCTT fnuDII/mvnI fnuDII/mvnI bsh1236I hhaI/cfoI ACCGTCTCCG GGAGCTGCAT GTGTCAGAGG TTTTCACCGT CATCACCGAA ACGCGCGAGG thal muli bsh1236I bstUI fnuDII/mvnI hinPI thaI bstul bsh1236I acil bstUI thaI nlaIII GAAGACGAAA GGGCCICGIG AIACGCCIAI IITIAIAGGI IAAIGICAIG **DSPHI** CTTCTGCTTT CCCGGAGCAC TATGCGGATA AAAATATCCA ATTACAGTAC rcaI hphI tru9I mseI hphI mnlI ahaII/bsaHI hinII/acvI caull alul nlaIII eco01091/draII haeIII/palI nspHI nspI fnu4HI mnlI bbvI sau96I asuI hpall scrFI Idsm nciI dsaV bsmAI esp3I bslI Ilodm bpuAI Isqq 2901 2831

bsmAI

rcal

bsrBI nlaIII

TCATGAGACA ATAACCCTGA GCCTTGGGGA TAAACAAATA AAAAGATTTA TGTAAGTTTA TACATAGGCG AGTACTCTGT TATTGGGACT acil bspHI ATTTGTTTAT TTTTCTAAAT ACATTCAAAT ATGTATCCGC

CGGAACCCCT

3001

nlaIV

earI/ksp632I Ilodm

TAAATGCTTC AATAATATTG AAAAAGGAAG AGTATGAGTA TTCAACATTT CCGTGTGGCC CTTATTCCCT

ATTTACGAAG TTATTATAAC TTTTTCCTTC TCATACTCAT AAGTTGTAAA GGCACAGGGG GAATAAGGGA

3071

sfaNI hphI hphI fnu4HI acil

CTCACCCAGA AACGCTGGTG AAAGTAAAAG TIGCGACCAC ITICATITIC GAGTGGGTCT TITITGCGGC ATTITGCCTI CCTGTTTTTG AAAAACGCCG TAAAACGGAA GGACAAAAAC 3141

hgiAI/aspHI bsp1286 **bsiHKAI** 

mpoI/ndeII[dam-]

sau3AI

dpnII[dam-] bstYI/xhoII

dpnI[dam+]

mboI/ndeII[dam-] dpnI[dam+] bmyI sau3AI

dpnII[dam-]

aciI

alwI[dam-]

taqI

bsrI

nspBII

apaLI/snoI mpoII[dam-]

GTTACATCGA ACTGGATCTC AACAGCGGTA maeIII ATGCTGAAGA TCAGTTGGGT GCACGAGTGG alw441/snoI eco57I

TGACCTAGAG TTGTCGCCAT CAATGTAGCT TACGACTTCT AGTCAACCCA CGTGCTCACC

FIG.32T

					iraI	LTC	<b>AAG</b>
		ipHI	tru9I	mseI	bmyI ahaIII/draI	TTTAAAG	AAATTTCA
		hgiAI/aspHI	bsp1286 tru9I	bsiHKAI mseI	DmyI	CTIGA GAGITITCGC CCCGAAGAAC GITIICCAAI GAIGAGCACI IITAAAGIIC	TCTAGGAACT CTCAAAAGCG GGGCTTCTTG CAAAAGGTTA CTACTCGTGA AAATTTCAAG
	maeII	psp1406I		00		GTTTTCCAAT	CAAAAGGTTA
	mae	dsd	xmnI	asp700	Ilodm	CCCGAAGAAC	GGGCTTCTTG
	/ndeII[dam-]		·			GAGTTTTCGC	CTCAAAAGCG
SausAi	mbol/ndeII	dpnI[dam+]	dpnII[dam-	.alwI[dam-]	bstYI/xhoII	AGATCCTTGA	TCTAGGAACT
				•		3261	

;					acil	bcgI mcrI fnu4HI	AGAGCAACTC GGTCGCCGCA	TCTCGTTGAG CCAGCGGCGT
scrFI nciI	Idsm	hpali	dsaV	hinl1/acy1	hgal caull	ahaII/bsaHI	TCCCGTGATG ACGCCGGGCA	AGGGCACTAC TGCGGCCCGT
acil	thaI	fnuDII/mvnI	bstUI	bsh1236I	hinPI	hhaI/cfoI	3321 TGCTATGTGG CGCGGTATTA TCCCGTGATG ACGCCGGGCA A	ACGATACACC GCGCCATAAT AGGGCACTAC TGCGGCCCGT TCTCGTTGAG CCAGCGGCGT

fokI 3381 TACACTATTC TCAGAATGAC TTGGTTGAGT ACTCACCAGT CACAGAAAAG CATCTTACGG ATGTGATAAG AGTCTTACTG AACCAACTCA TGAGTGGTCA GTGTCTTTTC GTAGAATGCC sfaNI scal hphl maelll bsrI csp61 rsal ddeI

FIG.32U

fnu4HI

bbvI

TACCGTACTG TCATTCTCTT AATACGTCAC GACGGTATTG GTACTCACTA TTGTGACGCC GGTTGAATGA CATGAGTGAT AACACTGCGG CCAACTTACT haeIII/palI eaeI cfrI fnu4HI aciI nlaIII 3441 ATGGCATGAC AGTAAGAGAA TTATGCAGTG CTGCCATAAC fnu4HI bbvI nlaIII

mboI/ndeII[dam-] sau3AI maeIII dpnII[damdpnI[dam+] nlaIII mboI/ndeII[dam-] avaII asuI dpnII[dam-] dpnI[dam+]

sau96I

sau3AI

AGACTGTTGC TAGCCTCCTG GCTTCCTCGA TTGGCGAAAA AACGTGTTGT ACCCCCTAGT ACATTGAGCG ICTGACAACG ATCGGAGGAC CGAAGGAGCT AACCGCTTTT TTGCACAACA TGGGGGATCA TGTAACTCGC nlaIII aciI aluI mnll mcrI 3511

alwI[dam-]

mbol/ndeII[dam-] aluI hpall mspI nlaIV dpnI[dam+] sau3AI

CGACTTACTT CGGTATGGTT TGCTGCTCGC ACTGTGGTGC TACGGTCGTC GCTGAATGAA GCCATACCAA ACGACGAGCG TGACACCACG ATGCCAGCAG sfaNI maelii CTTGATCGTT GGGAACCGGA GAACTAGCAA CCCTTGGCCT dpnII[dam-] bsaWI 3581

FIG.32V

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pvuI/bspCI

fnu4HI

fnuDII/mvnI bstUI

thaI

bbvI

bsh1236I

bsaI

bsmAI aciI

nlaIV hphI

lmdq/Ins6

CGACTATTTA

GACCAAATAA

3781 CTGGTTTATT GCTGATAAAT

mspl hpall cfr101 CTGGAGCCGG TGAGCGTGGG TCTCGCGGTA TCATTGCAGC GACCTCGGCC ACTCGCACCC AGAGCGCCAT AGTAACGTCG

mspI hpaII scrFI	aviII/fspl bsrI  maeII tru9I  psp1406I  3651 CAATGGCAAC AACTATTAA CTGGCGAACT ACTTACTCTA GCTCCGGC GTTACCGTTG TTGCAACGG TTTGATAATT GACCGCTTGA TGAATGAGAT CGAAGGGCCG	bgll sau961 sau961 haelII/pall msel bsrl acil avall hinPl asul mspl asel/asnl/vspl mnl1 asul hhal/cfol hpall 3711 AACAATTAAT AGACTGGATG GAGGCGGATA TTCAACGTCC TGGTGAAGAC GCGAGCCGG AAGGCCGACC TTGTTAATTA TCTGACCTAC CTCCGCCTAT TTCAACGTCC TGGTGAAGAC GCGAGCCGGG AAGGCCGACC
	SGCGAACT AC	sau96I avaII asuI STTGCAGG ACC
	bsrI tru9I mseI TATTAA CTC	aciI I GCGGATA AAC
l /cfoI	/fspI	ac mnll GAGGC
hinPI hhaI/cfoI mstI	avill/fspl maell psp14061 AACGTTGCGC AAAC TTGCAACGCG TTTG	tru9I fokI mseI bsrI aseI/asnI/vspI ATTAAT AGACTGGATG
	CAATGGCAAC GTTACCGTTG	tru9I mseI aseI/a: AACAATTAAT
	3651	3711

FIG.32W

maeII

nlallI

tru9I mseI

GGAGGGCATA GCATCAATAG ATGTGCTGCC CCTCAGTCCG CCTCCCGTAT CGTAGTTATC TACACGACGG GGAGTCAGGC hinfI pleI eam1105I mnlI TGACCCCGGT CTACCATTCG ACTGGGGCCA GATGGTAAGC bsrI haeIII/palI sau96I nlaIV asuI

tru9I nlaIV

ddeI

sau3AI

hgiCI mbol/ndeIl[dam-] dpnI[dam+]

mseI GACAGATCGC TGAGATAGGT GCCTCACTGA banI mnlI dpnII[dam-]

fokI

3901

TTAAGCATTG CTGTCTAGCG ACTCTATCCA CGGAGTGACT AATTCGTAAC TTGATACCTA CTTGCTTTAT GAACGAAATA AACTATGGAT

tru9I

tru9I mseI GTAACTGTCA GACCAAGTTT ACTCATATAT ACTTTAGATT GATTTAAAAC TTCATTTTTA TGAGTATATA TGAAATCTAA CTAAATTTTG AAGTAAAAAT ahaIII/draI mseI maeIII 3961

mbol/ndell[dam-] sau3AI sau3AI hphI rmaI

CTGGTTCAAA

CATTGACAGT

dpnI[dam+] mbol/ndeII[dam-] dpnI[dam+]

dpnII[dam-] alwI[dam-] dpnII[dam-] bstYI/xhoII tru9I

DspHI rcal bstYI/xhoII ahalii/drai maei .mboli[dam-] alwI[dam-] mseI

ATTTAAAAGG ATCTAGGTGA AGATCCTTTT TGATAATCTC ATGACCAAAA TCCCTTAACG TGAGTTTTCG IAAATTTTCC TAGATCCACT TCTAGGAAAA ACTATTAGAG TACTGGTTTT AGGGAATTGC ACTCAAAAGC

```
CCAAATACTG
                                                                                                                                                                                                                                                                                                                                                                                                                                                             GGTTTATGAC
                                               mboI/ndeII[dam-
                                                                                                                                                                                                                                                                                                                                                                                                                                               GCTTCAGCAG AGCGCAGATA
                                                                                                                                                                                                                                                                                                                                                                                                                                                              TCGCGTCTAT
                                                                                                                                                        AAGGTGACTC GCAGTCTGGG GCATCTTTTC TAGTTTCCTA GAAGAACTCT AGGAAAAAA
                                                                                                                                                                                                                                                                                    GGTTTGTTTG
                                                                                                                                                                                                                                                                                                   CCAAACAAAC
                                                                                                                                         4091 TICCACIGAG CGICAGACCC CGIAGAAAG AICAAAGGAI CIICIIGAGA ICCITIIIII
                                                                                                                                                                                                                                                                                                                                                                                                                                 hhaI/cfoI
                                                                           dpnII[dam-
                                                                                                                          bstYI/xhoII
                                                                                            alwI[dam-]
                                                             dpnI[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                                  hinPI
                               sau3AI
                mboI/ndeII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                              CGAAGTCGTC
                                                                                                                                                                                                                                                                                    GCAAACAAAA AAACCACCGC TACCAGCGGT
                                                                                                                                                                                                                                                                                                    TTTGGTGGCG ATGGTCGCCA
                                                                                                                                                                                                                                                     acil
                                                                                                                                                                                                                                                                     nspBII
                                                                                                           dpnI[dam+] mboII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                               eco57I
                                                dpnII[dam-]
                               dpnI[dam+]
                                                              bstYI/xhoII
                                                                             alwI[dam-]
                                                                                            mboI/ndeII[dam-]
sau3AI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                CCGGATCAAG AGCTACCAAC TCTTTTCCG AAGGTAACTG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                TTCCATTGAC
                                                                                                                                                                                                                                                                       acil
                                                                                                                                                                                                                                                                                                                                                                                                                  bsrI
                                                                                                                                                                                                                                                                                                                                                                                                                                 maeIII
                                                                                                                           dpnII[dam-]
                                                                            sau3AI
                                                                                                                                                                                                                                                                                                     GACGCGCATT AGACGACGAA CGTTTGTTTT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                TCGATGGTTG AGAAAAGGC
                                                                                                                                                                                                                                                                                      CTGCGCGTAA TCTGCTGCTT
                                                                                                                                                                                                                                                        fnu4HI
                                                                                                                                                                                                                                                                                                                                                     mboI/ndeII[dam-]
                                                                                                                                                                                                                                                                        bbvI
                                                                                                                                                                                                                                                                                                                                                                                                                                 aluI
                                                                                                                                                                                                          fnuDII/mvnI
                                                                                                             hgaI
                                                                                                                                                                                                                                                                                                                                                                                    dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                                     dpnI[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                   alwI[dam-]
                                                                                                                                                                                                                                                                       hhaI/cfoI
                                                                                                                                                                                                                                        bsh1236I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                GGCCTAGTTC
                                                                                                                                                                                                                                                                                                                                      sau3AI
                                                                                                                                                                                                                          bstUI
                                                                                                                                                                                                                                                        hinPI
                                                                                                                                                                                           thaI
                                                                                                                                                                                                                                                                                                                                                                                                                                 hpall
                                                                                                                                                                                                                                                                                       4151
                                                                                                                                                                                                                                                                                                                                                                                                                                                 4211
```

rmai haeiii/Pall scfi acii mnli maei bsli hael 4281 TCCTTCTAGT GTAGCCGTAG TTAGGCCACC ACTTCAAGAA CTCTGTAGCA CCGCCTACAT ACCTCGCTCT AGGAAGATCA CATCGGCATC AATCCGGTGG TGAAGTTCTT GAGACATCGT GGCGGATGTA TGGAGCGAGA	scrFI nciI mspI hpaII dsaV pleI	)
scfl ac CTCTGTAGCA CC GAGACATCGT GC		TCGTGTCTTA CARCACAGAAT G
ACTTCAAGAA (		TGGCGATAAG
haeiii/paii haei TTAGGCCACC ACT AATCCGGTGG TGA	fnu4HI bbvI nu4HI	bvI bsrI CTGCTGCCAG GACGACGGTC
bslI GTAGCCGTAG CATCGCCATC	fnu alwNI bbv bsrI fnu4HI	aeIII TTACCAGTGG
rmal mael TCCTTCTAGT AGGAAGATCA		m GCTAATCCTG CGATTAGGAC
4281	·	4351

hgiAI/aspHI bsp1286 bsp1286 bsiHKAI bbvI mcrI bhvI mcrI hinPI aciI hhaI/cfoI AGGCGCA GCGGTCGGC TGAACGGGG GTTCGTGCAC ACAGCCCAGC TTGGAGCGAA TCCGCGT CGCCAGCCCC CAAGCACGTG TGTCGTTCTT
ATA
mspI hpaII bsaWI maeIII 4421 TAGTTACCGG

CGACCTACAC CGAACTGAGA TACCTACAGC GTGAGCATTG AGAAAGCGCC ACGCTTCCCG AAGGGAGAAA GCTGGATGTG GCTTGACTCT ATGGATGTCG CACTCGTAAC TCTTTCGCGG TGCGAAGGGC TTCCCTCTTT FIG.32Z-1

hhaI/cfoI hinPI

haeII

scfI

4491

ecoRII mvaI GAGCGCACGA GGGAGCTTCC AGGGGGAAAC CTCGCGTGCT CCCTCGAAGG TCCCCCTTTG aluI apyI[dcm+] ecoRII bsaJI SCLFI bstNI dsaV mvaI hinPI mnlI hhaI/cfoI CGGAACAGGA GCCTTGTCCT GCGGCAGGGT CCGCCTGTCC ATAGGCCATT CGCCGTCCCA fnu4HI aciI GGCGGACAGG TATCCGGTAA hpall bsaWI Idsm bslI aciI

CGGACCATAG AAATATCAGG ACAGCCCAAA GCGGTGGAGA CTGAACTCGC AGCTAAAAAC ACTACGAGCA TGATGCTCGT sfaNI GCCIGGIAIC ITIAIAGICC IGICGGGIII CGCCACCICI GACIIGAGCG ICGAIIIIIG taqI hqaI drdI mnlI apyI[dcm+] bstNI dsaV

haeIII/palI

fnu4HI

acil thaI bslI fnuDII/mvnI bstUI

4701 CAGGGGGGG GAGCCTATGG AAAAACGCCA GCAACGCGGC GAGCCTATGG AAAAACGCCA GCAACGCGC CTCGGATACC TTTTTGCGGT CGTTGCGCCG

71/85

4631

scrFI

4561

4741 CTITITACGG TICCIGGCCT TITGCIGGCC TITIGCICAC AIGITCITIC CIGCGITAIC CCCIGALICI GAAAAATGCC AAGGACCGGA AAACGACCGG AAAACGAGTG TACAAGAAAG GACGCAATAG GGGACTAAGA hinfI tfil fnu4HI aflIII nspHI Idsu nlaIII haeIII/palI haeIII/palI apy1[dcm+.] mval bsll nlaIV haeI ecoRII bstNI SCLFI dsaV

CACCTATTGG CATAATGGCG GAAACTCACT CGACTATGGC GAGCGGCGTC GGCTTGCTGG CTTTGAGTGA GCTGATACCG CTCGCCGCAG CCGAACGACC mcrI fnu4HI acil bsrBI acil aluI GTGGATAACC GTATTACCGC aciI 4811

bbvI

fnu4HI
bbvI pleI
hinPI hinfI
mhoII
hhal/cfoI

mnlI

CTICTCGCGG GTTATGCGTT TGGCGGAGAG GAAGAGCGCC CAATACGCAA ACCGCCTCTC GAGCGCAGCG AGTCAGTGAG CGAGGAAGCG GCTCCTTCGC CTCGCGTCGC TCAGTCACTC

FIG.322-3

thaI

aciI CCCGCGCGTT GGCCGATTCA TTAATCCAGC TGGCACGACA GGTTTCCCGA CTGGAAAGCG GGGGGGGAA CCGGCTAAGT AATTAGGTCG ACCGTGCTGT CCAAAGGGCT GACCTTTCGC bsrI tfil asel/asnl/vspl msel nspBII aluI IInad tru9I bsh1236I haeIII/palI hinfI fnuDII/mvnI eaeI cfrI fnuDII/mvnI bsh1236I hhaI/cfoI bstuI hinPI bstuI thaI aciI bslI 4931

GGCAGTGAGC GCAACGCAAT TAATGTGAGT TACCTCACTC ATTAGGCACC CCAGGCTTTA CACTTTATGC CCGICACICG CGIIGCGIIA AITACACICA AIGGAGIGAG IAAICCGIGG GGICCGAAAI banI bsaJI hgiCI aseI/asnI/vspI mnlI maeIII

GTGAAATACG

apyI[dcm+]

bstNI

nlaIV

tru9I mseI

hhaI/cfoI

4991

hinPI

dsaV

ecoRII

SCrFI

mvaI

nlaIII GGAAACAGCT ATGACCATGA AAGGCCGAGC ATACAACACA CCTTAACACT CGCCTATTGT TAAAGTGTGT CCTTTGTCGA TACTGGTACT aluI GGAATTGTGA GCGGATAACA ATTTCACACA aciI bsrBI TICCGGCTCG TATGTIGTGT hpall Idsm 5061

16.322-5

>length: 5141

TTACGAATTA A AATGCTTAAT T

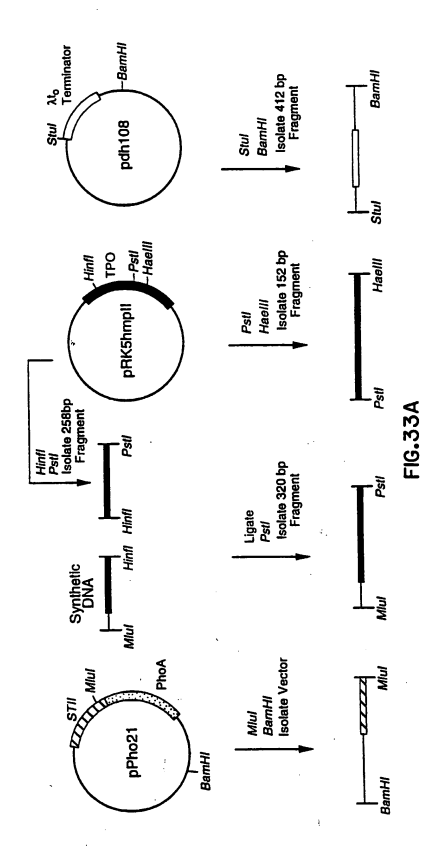
5131

xmnI asp700

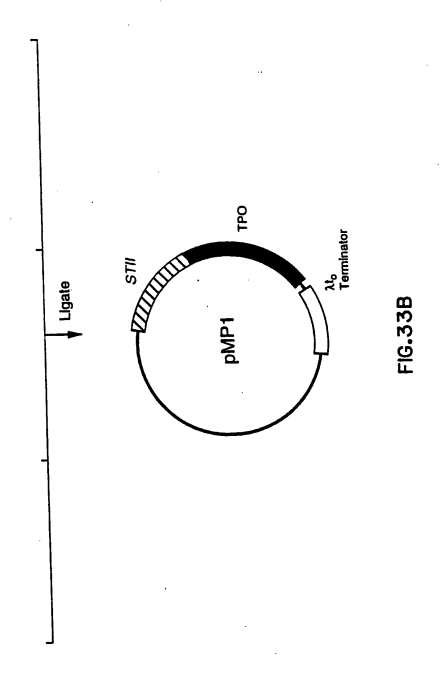
74/85 SUBSTITUTE SHEET (RULE 26)

aseI/asnI/vspI

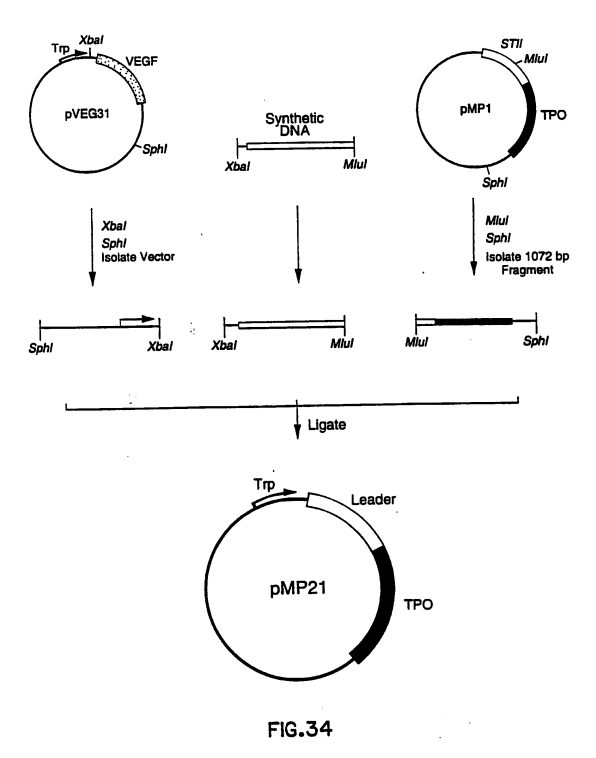
tru9I mseI



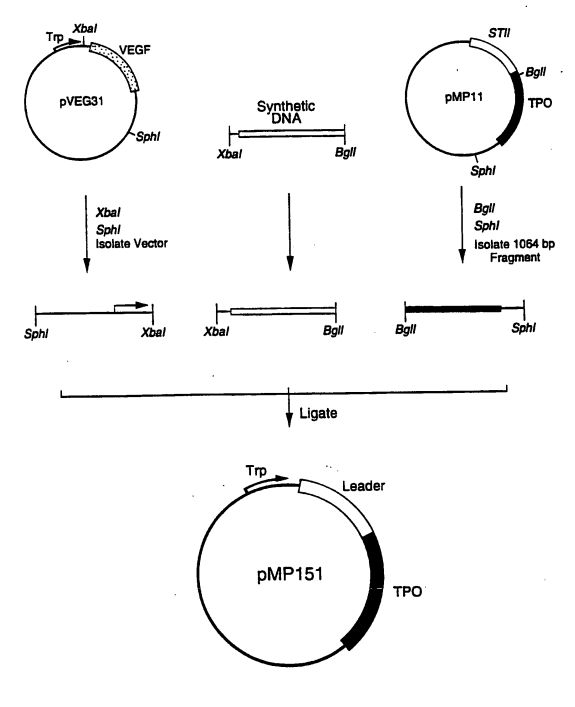
75/85



76/85 SUBSTITUTE SHEET (RULE 26)

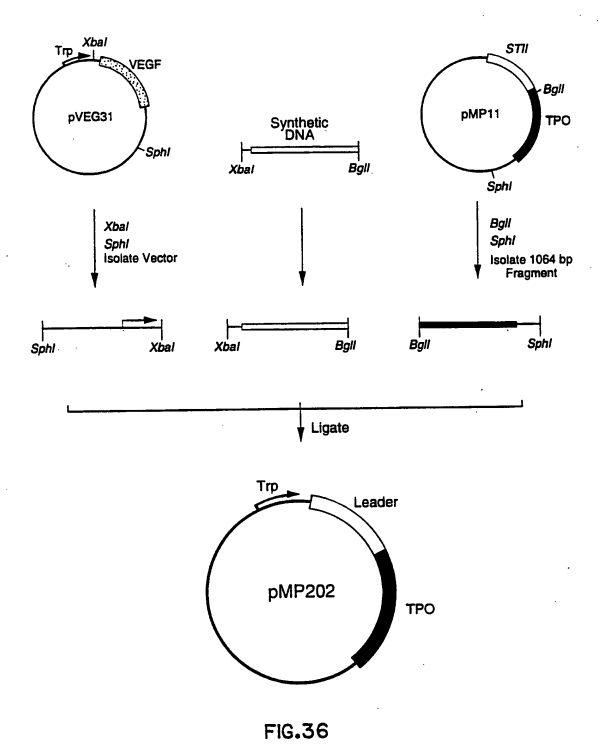


77/85

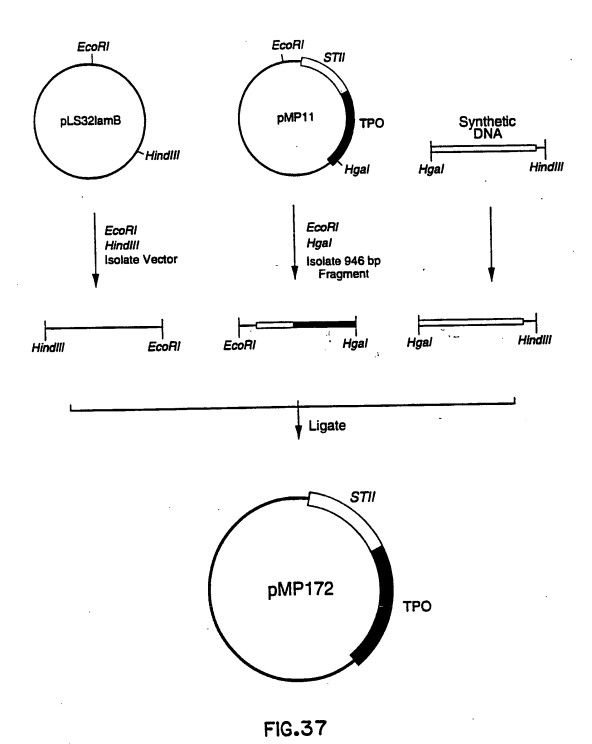


**FIG.35** 

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SUBSTITUTE SHEET (RULE 26)



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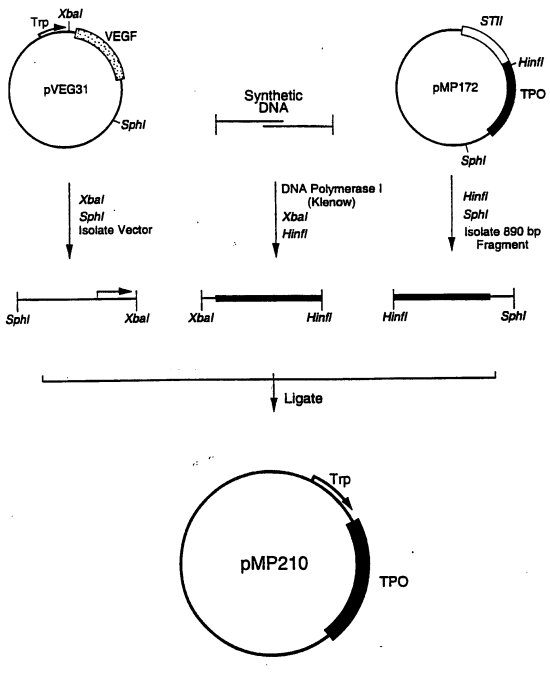


FIG.38

81/85 SUBSTITUTE SHEET (RULE 26)

	Met	Ser	Pro	Ala	Pro	Pro	Ala	
MP210 Bank	ATG	TCN	CCN	GCN	CCN	CCN	GCN	
MP210-1	ATG	TCT	CCA	GCG	CCG	CCA	GCG	
MP210-T8	ATG	TCG	CCT	GCT	CCA	CCT	GCT	
MP210-21	ATG	TCG	CCA	GCG	CCA	CCA	GCC	
MP210-24	ATG	TCC	CCA	GCC	CCA	CCC	GCA	
MP210-25	ATG	TĊG	CCA	GCG	CCG	CCA	GCG	

FIG.39

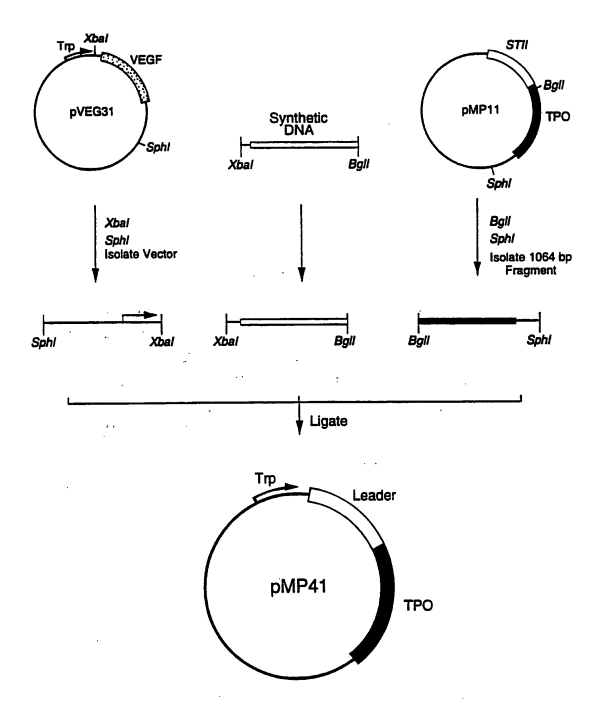


FIG.40

83/85 SUBSTITUTE SHEET (RULE 26)

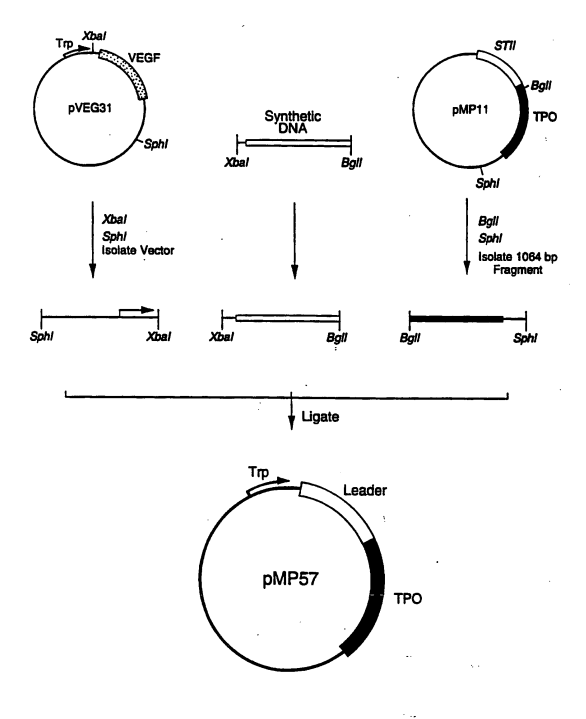


FIG.4 1

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SUBSTITUTE SHEET (RULE 26)

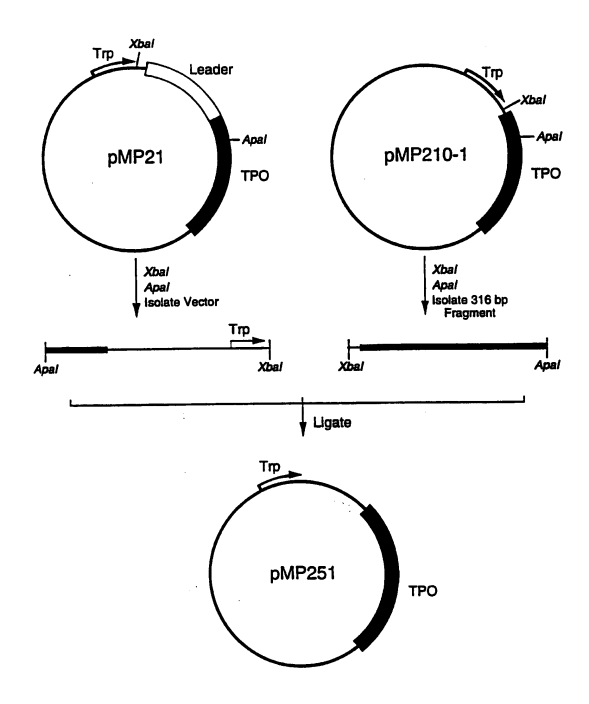


FIG.42

85/85 SUBSTITUTE SHEET (RULE 26)

## INTERNATIONAL SEARCH REPORT

Interna: 11 Application No
PCT/US 94/14553

	CONTION OF SURJECT MATTER		
IPC 6	FICATION OF SUBJECT MATTER C12N15/19 C07K14/52 C07K16/24	A61K38/19	
According to	International Patent Classification (IPC) or to both national classifica	tion and IPC	
B. FIELDS	SEARCHED  commentation searched (classification system followed by classification	symbols)	
Minimum do	CO7K		
<b>.</b> .			
Documentati	ion searched other than minimum documentation to the extent that suc	h documents are included in the fields sea	ırched
		,	
	ata base consulted during the international search (name of data base a	and, where practical, search terms used)	
Electrome a	ate pase commen aming are menuncial		
		•	
C. DOCUM	IENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.	
Category *	Citation of document, with indication, where appropriate, of the rele	vant passages	
	EXPERIMENTAL HEMATOLOGY,		1-5,12,
X	vol. 16, no. 3, March 1988		36-39
1	pages 201-205.		ì
	MCDONALD 'THROMBOPOETIN:ITS BIOLOGY, PURIFICATION, AND CHARACTES	RIZATION'	
	see the whole document		
			1-5,12,
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## INTERNATIONAL SEARCH REPORT

Intern val Application No
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